(12)

EUROPEAN PATENT SPECIFICATION

- (45) Date of publication and mention of the grant of the patent: 07.01.1999 Bulletin 1999/01
- (21) Application number: 92914973.0
- (22) Date of filing: 25.06.1992

- (51) Int Cl.6: **C12N 15/12**, C12P 21/02
- (86) International application number: PCT/US92/05374

(11)

(87) International publication number: WO 93/00432 (07.01.1993 Gazette 1993/02)

(54) BMP-9 COMPOSITIONS

BMP-9 ZUSAMMENSETZUNGEN COMPOSITIONS BMP-9

- (84) Designated Contracting States:

 AT BE CH DE DK ES FR GB GR IT LI LU MC NL SE
- (30) Priority: 25.06.1991 US 720590
- (43) Date of publication of application: 20.04.1994 Bulletin 1994/16
- (73) Proprietor: GENETICS INSTITUTE, INC. Cambridge, Massachusetts 02140 (US)
- (72) Inventors:
 - WOZNEY, John, M. Hudson, MA 01749 (US)
 - CELESTE, Anthony, J. Hudson, MA 01749 (US)

- (74) Representative: Jaenichen, Hans-Rainer, Dr. Vossius & Partner, Postfach 86 07 67 81634 München (DE)
- (56) References cited: WO-A-90/11366

WO-A-91/18098

PROCEEDINGS OF THE NATIONAL ACADEMY
OF SCIENCES OF USA vol. 87, no. 24, December
1990, WASHINGTON US pages 9843 - 9847
CELESTE, A.J. ET AL. 'Identification of
transforming growth factor beta family members
present in bone-inductive protein purified from
bovine bone'

P 0 592 562 B1

Note: Within nine months from the publication of the mention of the grant of the European patent, any person may give notice to the European Patent Office of opposition to the European patent granted. Notice of opposition shall be filed in a written reasoned statement. It shall not be deemed to have been filed until the opposition fee has been paid. (Art. 99(1) European Patent Convention).

Description

5

10

15

20

25

30

40

45

50

55

The present invention relates to a novel family of purified proteins designated BMP-9 proteins and processes for obtaining them. These proteins may be used to induce bone and/or cartilage formation and in wound healing and tissue repair.

The murine BMP-9 DNA sequence (SEQ ID NO: 1) and amino acid sequence (SEQ ID NO: 2) are set forth in Figure 1. Human BMP-9 sequence is set forth in Figure 3 (SEQ ID NO: 8 and SEQ ID NO: 9). It is contemplated that BMP-9 proteins are capable of inducing the formation of cartilage and/or bone. BMP-9 proteins may be further characterized by the ability to demonstrate cartilage and/or bone formation activity in the rat bone formation assay described below

Murine BMP-9 is characterized by comprising amino acid #319 to #428 of Figure 1 (SEQ ID NO: 2 amino acids #1-110). Murine BMP-9 may be produced by culturing a cell transformed with a DNA sequence comprising nucleotide #610 to nucleotide #1893 as shown in Figure 1 (SEQ ID NO: 1) and recovering and purifying from the culture medium a protein characterized by the amino acid sequence comprising amino acid #319 to #428 as shown in Figure 1 (SEQ ID NO: 2) substantially free from other proteinaceous materials with which it is co-produced.

Human BMP-9 is expected to be homologous to murine BMP-9 and is characterized by comprising amino acid #1 (Ser, Ala, Gly) to #110 of Figure 3 (SEQ ID NO: 9) (Arg). The invention includes methods for obtaining the DNA sequences encoding human BMP-9. This method entails utilizing the murine BMP-9 nucleotide sequence or portions thereof to design probes to screen libraries for the human gene or fragments thereof using standard techniques. Human BMP-9 may be produced by culturing a cell transformed with the BMP-9 DNA sequence and recovering and purifying BMP-9 from the culture medium. The expressed protein is isolated, recovered, and purified from the culture medium. The purified expressed protein is substantially free from other proteinaceous materials with which it is co-produced, as well as from other contaminants. The recovered purified protein is contemplated to exhibit cartilage and/or bone formation activity. The proteins of the invention may be further characterized by the ability to demonstrate cartilage and/or bone formation activity in the rat bone formation assay described below.

Human BMP-9 may be produced by culturing a cell transformed with a DNA sequence comprising nucleotide #124 to #453 as shown in SEQ ID NO: 8 and recovering and purifying from the culture medium a protein characterized by the amino acid sequence of SEQ ID NO: 9 from amino acid #1 to amino acid #110 substantially free from other proteinaceous materials with which it is co-produced.

Another aspect of the invention provides pharmaceutical compositions containing a therapeutically effective amount of a BMP-9 protein in a pharmaceutically acceptable vehicle or carrier. BMP-9 compositions of the invention may be used in the formation of cartilage. These compositions may further be utilized for the formation of bone. BMP-9 compositions may also be used for wound healing and tissue repair. Compositions of the invention may further include at least one other therapeutically useful agent such as the BMP proteins BMP-1, BMP-2, BMP-3, BMP-4, BMP-5, BMP-6, and BMP-7 disclosed for instance in PCT publications W088/00205, W089/10409, and W090/11366, and BMP-8, disclosed in U.S. application Ser. No. 07/641,204 filed January 15, 1991, Ser. No. 07/525,357 filed May 16, 1990, and Ser. No. 07/800,364 filed November 20, 1991.

The compositions of the invention may comprise, in addition to a BMP-9 protein, other therapeutically useful agents including growth factors such as epidermal growth factor (EGF), fibroblast growth factor (FGF), transforming growth factor (TGF- α and TGF- β), and insulin-like growth factor (IGF). The compositions may also include an appropriate matrix for instance, for supporting the composition and providing a surface for bone and/or cartilage growth. The matrix may provide slow release of the osteoinductive protein and/or the appropriate environment for presentation thereof.

The BMP-9 compositions may be employed in methods for treating a number of bone and/or cartilage defects, periodontal disease and various types of wounds. These methods, according to the invention, entail administering to a patient needing such bone and/or cartilage formation wound healing or tissue repair, an effective amount of a BMP-9 protein. These methods may also entail the administration of a protein of the invention in conjunction with at least one of the novel BMP proteins disclosed in the co-owned applications described above. In addition, these methods may also include the administration of a BMP-9 protein with other growth factors including EGF, FGF, TGF-α, TGF-β, and IGF.

Still a further aspect of the invention are DNA sequences coding for expression of a BMP-9, protein. Such sequences include the sequence of nucleotides in a 5' to 3' direction illustrated in Figure 1 (SEQ ID NO: 1) and Figure 3 (SEQ ID NO: 8) or DNA sequences which hybridize under stringent conditions with the DNA sequences of Figure 1 or 3 and encode a protein having the ability to induce the formation of cartilage and/or bone. Finally, allelic or other variations of the sequences of Figure 1 or 3, whether such nucleotide changes result in changes in the peptide sequence or not, are also included in the present invention.

A further aspect of the invention includes vectors comprising a DNA sequence as described above in operative association with an expression control sequence therefor. These vectors may be employed in a novel process for producing a BMP-9 protein of the invention in which a cell line transformed with a DNA sequence encoding a BMP-9

protein in operative association with an expression control sequence therefor, is cultured in a suitable culture medium and a BMP-9 protein is recovered and purified therefrom. This process may employ a number of known cells both prokaryotic and eukaryotic as host cells for expression of the polypeptide.

Other aspects and advantages of the present invention will be apparent upon consideration of the following detailed description and preferred embodiments thereof.

Brief Description of the Drawings

10

15

20

25

30

35

45

50

55

FIG. 1 comprises DNA sequence and derived amino acid sequence of murine BMP-9 from clone ML14a further described below.

FIG. 2 comprises DNA sequence and derived amino acid sequence of human BMP-4 from lambda U20S-3 ATCC #40342.

FIG. 3 comprises DNA sequence and derived amino acid sequence of human BMP-9 from λ FIX/H6III ATCC # 75252.

Detailed Descripton of the Invention

The murine BMP-9 nucleotide sequence (SEQ ID NO: 1) and encoded amino acid sequence (SEQ ID NO: 2) are depicted in Figure 1. Purified murine BMP-9 proteins of the present invention are produced by culturing a host cell transformed with a DNA sequence comprising the DNA coding sequence of Figure 1 (SEQ ID NO: 1) from nucleotide #610 to nucleotide #1893 and recovering and purifying from the culture medium a protein which contains the amino acid sequence or a substantially homologous sequence as represented by amino acid #319 to #428 of Figure 1 (SEQ ID NO: 2). The BMP-9 proteins recovered from the culture medium are purified by isolating them from other proteinaceous materials from which they are co-produced and from other contaminants present.

Human BMP-9 nucleotide and amino acid sequence is depicted in SEQ ID No: 8 and 9. Mature human BMP-9 is expected to comprise amino acid #1 (Ser, Ala, Gly) to #110 (Arg).

Human BMP-9 may be produced by culturing a cell transformed with a DNA sequence comprising nucleotide #124 to #453 as shown in SEQ ID NO: 8 and recovering and purifying from the culture medium a protein characterized by the amino acid sequence of SEQ ID NO: 9 from amino acid #1 to amino acid #110 substantially free from other proteinaceous materials with which it is co-produced.

BMP-9 proteins may be characterized by the ability to induce the formation of cartilage. BMP-9 proteins may be further characterized by the ability to induce the formation of bone. BMP-9 proteins may be further characterized by the ability to demonstrate cartilage and/or bone formation activity in the rat bone formation assay described below.

The BMP-9 proteins provided herein also include factors encoded by sequences similar to those of Figure 1 and 3 (SEQ ID NO's: 1 and 8), but into which modifications are naturally provided (e.g. allelic variations in the nucleotide sequence which may result in amino acid changes in the polypeptide) or deliberately engineered. For example, synthetic polypeptides may wholly or partially duplicate continuous sequences of the amino acid residues of Figure 1 of Figure 3 (SEQ ID NO's: 2 and 9). These sequences, by virtue of sharing primary, secondary, or tertiary structural and conformational characteristics with bone growth factor polypeptides of Figure 1 and Figure 3 may possess bone growth factor biological properties in common therewith. Thus, they may be employed as biologically active substitutes for naturally-occurring BMP-9 and other BMP-9 polypeptides in therapeutic processes.

Other specific mutations of the sequences of BMP-9 proteins described herein involve modifications of glycosylation sites. These modifications may involve O-linked or N-linked glycosylation sites. For instance, the absence of glycosylation or only partial glycosylation results from amino acid substitution or deletion at asparagine-linked glycosylation recognition sites. The asparagine-linked glycosylation recognition sites comprise tripeptide sequences which are specifically recognized by appropriate cellular glycosylation enzymes. These tripeptide sequences are either asparagine-X-threonine or asparagine-X-serine, where X is usually any amino acid. A variety of amino acid substitutions or deletions at one or both of the first or third amino acid positions of a glycosylation recognition site (and/or amino acid deletion at the second position) results in non-glycosylation at the modified tripeptide sequence.

The present invention also encompasses the novel DNA sequences, free of association with DNA sequences encoding other proteinaceous materials, and coding on expression for BMP-9 proteins. These DNA sequences include those depicted in Figure 1 or Figure 3 (SEQ ID NO's: 1 and 8) in a 5' to 3' direction and those sequences which hybridize thereto under stringent hybridization conditions [see, T. Maniatis et al, Molecular Cloning (A Laboratory Manual), Cold Spring Harbor Laboratory (1982), pages 387 to 389] and encode a protein having cartilage and/or bone inducing activity.

Similarly, DNA sequences which code for BMP-9 proteins coded for by the sequences of Figure 1 or Figure 3, but which differ in codon sequence due to the degeneracies of the genetic code or allelic variations (naturally-occurring base changes in the species population which may or may not result in an amino acid change) also encode the novel factors described herein. Variations in the DNA sequences of Figure 1 or Figure 3 (SEQ ID NO: 1 and 8) which are

caused by point mutations or by induced modifications (including insertion, deletion, and substitution) to enhance the activity, half-life or production of the polypeptides encoded are also encompassed in the invention.

Another aspect of the present invention provides a novel method for producing BMP-9 proteins. The method of the present invention involves culturing a suitable cell line, which has been transformed with a DNA sequence encoding a BMP-9 protein of the invention, under the control of known regulatory sequences. The transformed host cells are cultured and the BMP-9 proteins recovered and purified from the culture medium. The purified proteins are substantially free from other proteins with which they are co-produced as well as from other contaminants.

5

10

15

20

25

30

35

40

45

50

Suitable cells or cell lines may be mammalian cells, such as Chinese hamster ovary cells (CHO). The selection of suitable mammalian host cells and methods for transformation, culture, amplification, screening, product production and purification are known in the art. See, e.g., Gething and Sambrook, Nature, 293:620-625 (1981), or alternatively, Kaufman et al, Mol. Cell. Biol., 5(7):1750-1759 (1985) or Howley et al, U.S. Patent 4,419,446. Another suitable mammalian cell line, which is described in the accompanying examples, is the monkey COS-1 cell line. The mammalian cell CV-1 may also be suitable.

Bacterial cells may also be suitable hosts. For example, the various strains of <u>E. coli</u> (e.g., HB101, MCl061) are well-known as host cells in the field of biotechnology. Various strains of <u>B. subtilis, Pseudomonas</u>, other bacilli and the like may also be employed in this method.

Many strains of yeast cells known to those skilled in the art may also be available as host cells for expression of the polypeptides of the present invention. Additionally, where desired, insect cells may be utilized as host cells in the method of the present invention. See, e.g. Miller et al, <u>Genetic Engineering</u>, <u>8</u>:277-298 (Plenum Press 1986) and references cited therein.

Another aspect of the present invention provides vectors for use in the method of expression of these novel BMP-9 polypeptides. Preferably the vectors contain the full novel DNA sequences described above which encode the novel factors of the invention. Additionally the vectors also contain appropriate expression control sequences permitting expression of the BMP-9 protein sequences. Alternatively, vectors incorporating modified sequences as described above are also embodiments of the present invention. The vectors may be employed in the method of transforming cell lines and contain selected regulatory sequences in operative association with the DNA coding sequences of the invention which are capable of directing the replication and expression thereof in selected host cells. Regulatory sequences for such vectors are known to those skilled in the art and may be selected depending upon the host cells. Such selection is routine and does not form part of the present invention.

A protein of the present invention, which induces cartilage and/or bone formation in circumstances where bone is not normally formed, has application in the healing of bone fractures and cartilage defects in humans and other animals. Such a preparation employing a BMP-9 protein may have prophylactic use in closed as well as open fracture reduction and also in the improved fixation of artificial joints. De novo bone formation induced by an osteogenic agent contributes to the repair of congenital, trauma induced, or oncologic resection induced craniofacial defects, and also is useful in cosmetic plastic surgery. A BMP-9 protein may be used in the treatment of periodontal disease, and in other tooth repair processes. Such agents may provide an environment to attract bone-forming cells, stimulate growth of bone-forming cells or induce differentiation of progenitors of bone-forming cells. BMP-9 polypeptides of the invention may also be useful in the treatment of osteoporosis. A variety of osteogenic, cartilage-inducing and bone inducing factors have been described. See, e.g. European patent applications 148,155 and 169,016 for discussions thereof.

The proteins of the invention may also be used in wound healing and related tissue repair. The types of wounds include, but are not limited to burns, incisions and ulcers. (See, e.g. PCT Publication W084/01106 for discussion of wound healing and related tissue repair).

It is further contemplated that proteins of the invention may increase neuronal survival and therefore be useful in transplantation and treatment of conditions exhibiting a decrease in neuronal survival.

A further aspect of the invention is a therapeutic method and composition for repairing fractures and other conditions related to cartilage and/or bone defects or periodontal diseases. The invention further comprises therapeutic methods and compositions for wound healing and tissue repair. Such compositions comprise a therapeutically effective amount of at least one of the BMP-9 proteins of the invention in admixture with a pharmaceutically acceptable vehicle, carrier or matrix.

It is expected that the proteins of the invention may act in concert with or perhaps synergistically with other related proteins and growth factors. Further therapeutic methods and compositions of the invention therefore comprise a therapeutic amount of at least one BMP-9 protein of the invention with a therapeutic amount of at least one of the other BMP proteins disclosed in co-owned applications described above. Such combinations may comprise separate molecules of the BMP proteins or heteromolecules comprised of different BMP moieties. For example, a method and composition of the invention may comprise a disulfide linked dimer comprising a BMP-9 protein subunit and a subunit from one of the "BMP" proteins described above. A further embodiment may comprise a heterodimer of BMP-9 moieties. Further, BMP-9 proteins may be combined with other agents beneficial to the treatment of the bone and/or cartilage defect, wound, or tissue in question. These agents include various growth factors such as epidermal growth factor

(EGF), platelet derived growth factor (PDGF), transforming growth factors (TGF- α and TGF- β), and insulin-like growth factor (IGF).

The preparation and formulation of such physiologically acceptable protein compositions, having due regard to pH, isotonicity, stability and the like, is within the skill of the art. The therapeutic compositions are also presently valuable for veterinary applications due to the lack of species specificity in BMP proteins. Particularly domestic animals and thoroughbred horses in addition to humans are desired patients for such treatment with BMP-9 of the present invention.

The therapeutic method includes administering the composition topically, systemically, or locally as an implant or device. When administered, the therapeutic composition for use in this invention is, of course, in a pyrogen-free, physiologically acceptable form. Further, the composition may desirably be encapsulated or injected in a viscous form for delivery to the site of bone, cartilage or tissue damage. Topical administration may be suitable for wound healing and tissue repair. Therapeutically useful agents other than the BMP-9 proteins which may also optionally be included in the composition as described above, may alternatively or additionally, be administered simultaneously or sequentially with the BMP composition in the methods of the invention.

Preferably for bone and/or cartilage formation, the composition would include a matrix capable of delivering BMP-9 or other BMP proteins to the site of bone and/or cartilage damage, providing a structure for the developing bone and cartilage and optimally capable of being resorbed into the body. The matrix may provide slow release of BMP-9 and/or the appropriate environment for presentation thereof. Such matrices may be formed of materials presently in use for other implanted medical applications.

The choice of matrix material is based on biocompatibility, biodegradability, mechanical properties, cosmetic appearance and interface properties. The particular application of the BMP-9 compositions will define the appropriate formulation. Potential matrices for the compositions may be biodegradable and chemically defined calcium sulfate, tricalciumphosphate, hydroxyapatite, polylactic acid and polyanhydrides. Other potential materials are biodegradable and biologically well defined, such as bone or dermal collagen. Further matrices are comprised of pure proteins or extracellular matrix components. Other potential matrices are nonbiodegradable and chemically defined, such as sintered hydroxyapatite, bioglass, aluminates, or other ceramics. Matrices may be comprised of combinations of any of the above mentioned types of material, such as polylactic acid and hydroxyapatite or collagen and tricalciumphosphate. The bioceramics may be altered in composition, such as in calcium-aluminate-phosphate and processing to alter pore size, particle size, particle shape, and biodegradability.

The dosage regimen will be determined by the attending physician considering various factors which modify the action of the BMP-9 protein, e.g. amount of bone weight desired to be formed, the site of bone damage, the condition of the damaged bone, the size of a wound, type of damaged tissue, the patient's age, sex, and diet, the severity of any infection, time of administration and other clinical factors. The dosage may vary with the type of matrix used in the reconstitution and the types of BMP proteins in the composition. The addition of other known growth factors, such as IGF I (insulin like growth factor I), to the final composition, may also effect the dosage. Progress can be monitored by periodic assessment of bone growth and/or repair, for example, x-rays, histomorphometric determinations and tetracycline labeling.

The following examples illustrate practice of the present invention in recovering and characterizing murine BMP-9 protein and employing it to recover the human and other BMP-9 proteins, obtaining the human proteins and expressing the proteins via recombinant techniques.

EXAMPLE I

5

10

15

20

25

30

35

40

45

50

55

Murine BMP-9

750,000 recombinants of a mouse liver cDNA library made in the vector lambdaZAP (Stratagene/Catalog #935302) are plated and duplicate nitrocellulose replicas made. A fragment of human BMP-4 DNA corresponding to nucleotides 1330-1627 of Figure 2 (SEQ ID NO: 3) (the human BMP-4 sequence) is ³²P-labeled by the random priming procedure of Feinberg et al. [Anal. Biochem. 132: 6-13 (1983)] and hybridized to both sets of filters in SHB at 60°C for 2 to 3 days. Both sets of filters are washed under reduced stringency conditions (4X SSC, 0.1% SDS at 60°C). Many duplicate hybridizing recombinants of various intensities (approximately 92) are noted. 50 of the strongest hybridizing recombinant bacteriophage are plaque purified and their inserts are transferred to the plasmid Bluescript SK (+/-) according to the in vivo excision protocol described by the manufacturer (Stratagene). DNA sequence analysis of several recombinants indicate that they encode a protein homologous to other BMP proteins and other proteins in the TGF-β family. The DNA sequence and derived amino acid sequence of one recombinant, designated ML14a, is set forth in Figure 1. (SEQ ID NO: 1)

The nucleotide sequence of clone ML14a contains an open reading frame of 1284 bp, encoding a BMP-9 protein of 428 amino acids. The encoded 428 amino acid BMP-9 protein is contemplated to be the primary translation product as the coding sequence is preceded by 609 bp of 5' untranslated sequence with stop codons in all three reading frames.

The 428 amino acid sequence predicts a BMP-9 protein with a molecular weight of 48,000 daltons.

Based on knowledge of other BMP proteins and other proteins within the TGF-β family, it is predicted that the precursor polypeptide would be cleaved at the multibasic sequence ARG-ARG-LYS-ARG in agreement with a proposed consensus proteolytic processing sequence of ARG-X-X-ARG. Cleavage of the BMP-9 precursor polypeptide at this location would generate a 110 amino acid mature peptide beginning with the amino acid SER at position #319. The processing of BMP-9 into the mature form is expected to involve dimerization and removal of the N-terminal region in a manner analogous to the processing of the related protein TGF-β [L.E. Gentry, et al., Molec. & Cell. Biol. 8:4162 (1988); R. Derynck, et al., Nature 316:701 (1985)].

It is contemplated therefore that the mature active species of murine BMP-9 comprises a homodimer of 2 polypeptide subunits, each subunit comprising amino acids #319-#428 with a predicted molecular weight of approximately 12,000 daltons. Further active species are contemplated comprising amino acids #326 - #428 thereby including the first conserved cysteine residue. As with other members of the BMP and TGF- β family of proteins, the carboxy-terminal region of the BMP-9 protein exhibits greater sequence conservation than the more amino-terminal portion. The percent amino acid identity of the murine BMP-9 protein in the cysteine-rich C-terminal domain (amino acids #326 - #428) to the corresponding region of other human BMP proteins and other proteins within the TGF- β family is as follows: BMP-2, 53%; BMP-3, 43%; BMP-4, 53%; BMP-5, 55%; BMP-6, 55%; BMP-7, 53%; Vgl, 50%; GDF-1, 43%; TGF- β 1, 32%; TGF- β 2, 34%; TGF- β 3, 34%; inhibin β (B), 34%; and inhibin β (A), 42%.

EXAMPLE II

20

25

35

5

10

Human BMP-9

Murine and human osteoinductive factor genes are presumed to be significantly homologous, therefore the murine coding sequence or a portion thereof is used as a probe to screen a human genomic library or as a probe to identify a human cell line or tissue which synthesizes the analogous human cartilage and/or bone protein. A human genomic library (Toole et al., supra) may be screened with such a probe, and presumptive positives isolated and DNA sequence obtained. Evidence that this recombinant encodes a portion of the human BMP-9 relies of the murine/human protein and gene structure homologies.

Once a recombinant bacteriophage containing DNA encoding portion of the human cartilage and/or bone inductive factor molecule is obtained, the human coding sequence can be used as a probe to identify a human cell line or tissue which synthesizes BMP-9. Alternatively, the murine coding sequence can be used as a probe to identify such human cell line or tissue. Briefly described, RNA is extracted from a selected cell or tissue source and either electrophoresed on a formaldehyde agarose gel and transferred to nitrocellulose, or reacted with formaldehyde and spotted on nitrocellulose directly. The nitrocellulose is then hybridized to a probe derived from a coding sequence of the murine or human BMP-9. mRNA is selected by oligo (dT) cellulose chromatography and cDNA is synthesized and cloned in lambda gt10 or lambda ZAP by established techniques (Toole et al., supra).

Additional methods known to those skilled in the art may be used to isolate the human and other species' BMP-9 proteins of the invention.

40 A. Isolation of Human BMP-9 DNA

One million recombinants of a human genomic library constructed in the vector λ FIX (Stratagene catalog # 944201) are plated and duplicate nitrocellulose replicas made. Two oligonucleotides probes designed on the basis of nucleotides #1665-#1704 and #1837-#1876 of the sequence set forth in Figure 1 (SEQ ID NO:1) are synthesized on an automated DNA synthesizer. The sequence of these two oligonucleotides is indicated below:

#1: CTATGAGTGTAAAGGGGGTTGCTTCTTCCCATTGGCTGAT

50

55

45

#2: GTGCCAACCCTCAAGTACCACTATGAGGGGATGAGTGTGG

These two oligonucleotide probes are radioactively labeled with γ^{32} P-ATP and each is hybridized to one set of the duplicate nitrocellulose replicas in SHB at 65°C and washed with 1X SSC, 0.1% SDS at 65°C. Three recombinants which hybridize to both oligonucleotide probes are noted. All three positively hybridizing recombinants are plaque purified, bacteriophage plate stocks are prepared and bacteriophage DNA is isolated from each. The oligonucleotide hybridizing regions of one of these recombinants, designated HGIII, is localized to a 1.2 kb Pst I/Xba I fragment. This fragment is subcloned into a plasmid vector (pGEM-3) and DNA sequence analysis is performed. HGIII was deposited

with the ATCC, 12301 Parklawn Drive, Rockville, Maryland USA on June 16, 1992 under the requirements of the Budapest Treaty and designated as ATCC # 75252. This subclone is designated pGEM-111. A portion of the DNA sequence of clone pGEM-111 is set forth in Figure 3 (SEQ ID NO:8/ HUMAN BMP-9 sequence). This sequence encodes the entire mature region of human BMP-9 and a portion of the propeptide. It should be noted that this sequence consists of preliminary data. Particularly, the propeptide region is subject to further analysis and characterization. For example, nucleotides #1 through #3 (TGA) encode a translational stop which may be incorrect due to the preliminary nature of the sequence. It is predicted that additional sequences present in both pGEM-111 (the 1.2 kb Pstl/Xbal fragment of HGIII subcloned into pGEM) and HGIII encode additional amino acids of the human BMP-9 propeptide region. Based on knowledge of other BMPs and other proteins within the TGF-β family, it is predicted that the precursor polypeptide would be cleaved at the multibasic sequence ARG-ARG-LYS-ARG (amino acids # -4 through # -1 of SEQUENCE ID NO:9) in agreement with a proposed consensus proteolytic processing sequence ARG-X-X-ARG. Cleavage of the human BMP-9 precursor polypeptide at this location would generate a 110 amino acid mature peptide beginning with the amino acid SER at position #1 of SEQUENCE ID NO:9 (encoded by nucleotides #124 through #126 of SEQUENCE ID NO:8). The processing of human BMP-9 into the mature form is expected to involve dimerization and removal of the N-terminal region in a manner analogous to the processing of the related protein TGF-β [L.E. Gentry, et al., Molec. & Cell. Biol. 8:4162 (1988); R. Derynck, et al., Nature 316:701 (1985)].

It is contemplated therefore that the mature active species of human BMP-9 comprises a homodimer of two polypeptide subunits, each subunit comprising amino acids #1 through #110 of SEQUENCE ID NO:9, with a predicted molecular weight of 12,000 daltons. Further active species are contemplated comprising amino acids #8 through #110 thereby including the first conserved cysteine residue. As with other members of the BMP and TGF- β family of proteins, the carboxy-terminal portion of the human BMP-9 sequence exhibits greater sequence conservation than the aminoterminal portion. the percent amino acid identity of the human BMP-9 protein in the cysteine-rich C-terminal domain (amino acids #8 through #110) to the corresponding region of other human BMP proteins and other proteins within the TGF- β family is as follows: BMP-2, 52%; BMP-3, 40%; BMP-4, 52%; BMP-5, 55%; BMP-6, 55%; BMP-7, 53%; murine BMP-9, 97%; Vgl, 50%; GDF-1, 44%; TGF- β 1, 32%; TGF- β 2, 32%; TGF- β 3, 32%; inhibin β (B), 35%; and inhibin β (A), 41%.

EXAMPLE III

10

20

30

35

40

50

Rosen Modified Sampath-Reddi Assay

A modified version of the rat bone formation assay described in Sampath and Reddi, <u>Proc. Natl. Acad. Sci. U.S. A.</u>, 80:6591-6595 (1983) is used to evaluate bone and/or cartilage activity of the BMP proteins. This modified assay is herein called the Rosen-modified Sampath-Reddi assay. The ethanol precipitation step of the Sampath-Reddi procedure is replaced by dialyzing (if the composition is a solution) or diafiltering (if the composition is a suspension) the fraction to be assayed against water. The solution or suspension is then redissolved in 0.1 % TFA, and the resulting solution added to 20mg of rat matrix. A mock rat matrix sample not treated with the protein serves as a control. This material is frozen and lyophilized and the resulting powder enclosed in #5 gelatin capsules. The capsules are implanted subcutaneously in the abdominal thoracic area of 21 - 49 day old male Long Evans rats. The implants are removed after 7 - 14 days. Half of each implant is used for alkaline phosphatase analysis [See, A. H. Reddi et al., <u>Proc. Natl Acad Sci.</u>, 69:1601 (1972)].

The other half of each implant is fixed and processed for histological analysis. 1µm glycolmethacrylate sections are stained with Von Kossa and acid fuschin to score the amount of induced bone and cartilage formation present in each implant. The terms +1 through +5 represent the area of each histological section of an implant occupied by new bone and/or cartilage cells and matrix. A score of +5 indicates that greater than 50% of the implant is new bone and/or cartilage produced as a direct result of protein in the implant. A score of +4, +3, +2 and +1 would indicate that greater than 40%, 30%, 20% and 10% respectively of the implant contains new cartilage and/or bone. In a modified scoring method, three non-adjacent sections are evaluated from each implant and averaged. "+/-" indicates tentative identification of cartilage or bone; "+1" indicates >10% of each section being new cartilage or bone; "+2", >25%; "+3", >50%; "+4", -75%; "+5", >80%. A "-" indicates that the implant is not recovered.

It is contemplated that the dose response nature of the BMP-9 containing samples of the matrix samples will demonstrate that the amount of bone and/or cartilage formed increases with the amount of BMP-9 in the sample. It is contemplated that the control samples will not result in any bone and/or cartilage formation.

As with other cartilage and/or bone inductive proteins such as the above-mentioned "BMP" proteins, the bone and/or cartilage formed is expected to be physically confined to the space occupied by the matrix. Samples are also analyzed by SDS gel electrophoresis and isoelectric focusing followed by autoradiography. The activity is correlated with the protein bands and pl. To estimate the purity of the protein in a particular fraction an extinction coefficient of 1 OD/mgcm is used as an estimate for protein and the protein is run on SDS PAGE followed by silver staining or radioiodination

and autoradiography.

EXAMPLE IV

5 Expression of BMP-9

In order to produce murine, human or other mammalian BMP-9 proteins, the DNA encoding it is transferred into an appropriate expression vector and introduced into mammalian cells or other preferred eukaryotic or prokaryotic hosts by conventional genetic engineering techniques. The preferred expression system for biologically active recombinant human BMP-9 is contemplated to be stably transformed mammalian cells.

One skilled in the art can construct mammalian expression vectors by employing the sequence of Figure 1 (SEQ ID NO: 1) or Figure 3 (SEQ ID NO: 8), or other DNA sequences encoding BMP-9 proteins or other modified sequences and known vectors, such as pCD [Okayama et al., Mol. Cell Biol., 2:161-170 (1982)], pJL3, pJL4 [Gough et al., EMBO J., 4:645-653 (1985)] and pMT2 CXM.

The mammalian expression vector pMT2 CXM is a derivative of p91023 (b) (Wong et al., Science <u>228</u>: 810-815, 1985) differing from the latter in that it contains the ampicillin resistance gene in place of the tetracycline resistance gene and further contains a Xhol site for insertion of cDNA clones. The functional elements of pMT2 CXM have been described (Kaufman, R.J., 1985, Proc. Natl. Acad. Sci. USA <u>82</u>:689-693) and include the adenovirus VA genes, the SV40 origin of replication including the 72 bp enhancer, the adenovirus major late promoter including a 5' splice site and the majority of the adenovirus tripartite leader sequence present on adenovirus late mRNAs, a 3' splice acceptor site, a DHFR insert, the SV40 early polyadenylation site (SV40), and pBR322 sequences needed for propagation in <u>E. coli.</u>

Plasmid pMT2 CXM is obtained by EcoRI digestion of pMT2-VWF, which has been deposited with the American Type Culture Collection (ATCC), Rockville, MD (USA) under accession number ATCC 67122. EcoRI digestion excises the cDNA insert present in pMT2-VWF, yielding pMT2 in linear form which can be ligated and used to transform <u>E. coli</u> HB 101 or DH-5 to ampicillin resistance. Plasmid pMT2 DNA can be prepared by conventional methods. pMT2 CXM is then constructed using loopout/in mutagenesis [Morinaga, et al., <u>Biotechnology 84:</u> 636 (1984). This removes bases 1075 to 1145 relative to the Hind III site near the SV40 origin of replication and enhancer sequences of pMT2. In addition it inserts the following sequence:

5' PO-CATGGGCAGCTCGAG-3' (SEQ ID NO: 5)

at nucleotide 1145. This sequence contains the recognition site for the restriction endonuclease Xho I. A derivative of pMT2CXM, termed pMT23, contains recognition sites for the restriction endonucleases Pstl, Eco RI, Sall and Xhol. Plasmid pMT2 CXM and pMT23 DNA may be prepared by conventional methods.

pEMC2bl derived from pMT21 may also be suitable in practice of the invention. pMT21 is derived from pMT2 which is derived from pMT2-VWF. As described above EcoRl digestion excises the cDNA insert present in pMT-VWF, yielding pMT2 in linear form which can be ligated and used to transform <u>E. Coli</u> HB 101 or DH-5 to ampicillin resistance. Plasmid pMT2 DNA can be prepared by conventional methods.

pMT21 is derived from pMT2 through the following two modifications. First, 76 bp of the 5' untranslated region of the DHFR cDNA including a stretch of 19 G residues from G/C tailing for cDNA cloning is deleted. In this process, a Xhol site is inserted to obtain the following sequence immediately

upstream from DHFR: 5' - CTGCAGGCGAGCCTGAATTCCTCGAGCCATCATG-3'
PstI Eco RI XhoI
(SEQ ID NO: 6)

Second, a unique Clal site is introduced by digestion with EcoRV and Xbal, treatment with Klenow fragment of DNA polymerase I, and ligation to a Clal linker (CATCGATG). This deletes a 250 bp segment from the adenovirus associated RNA (VAI) region but does not interfere with VAI RNA gene expression or function. pMT21 is digested with EcoRI and Xhol, and used to derive the vector pEMC2B1.

A portion of the EMCV leader is obtained from pMT2-ECAT1 [S.K. Jung, et al, <u>J. Virol</u> 63:1651-1660 (1989)] by digestion with Eco RI and PstI, resulting in a 2752 bp fragment. This fragment is digested with TaqI yielding an Eco RI-TaqI fragment of 508 bp which is purified by electrophoresis on low melting agarose gel. A 68 bp adapter and its complementary strand are synthesized with a 5' TaqI protruding end and a 3' XhoI protruding end which has the following sequence:

35

40

45

50

10

15

5'-<u>CGA</u>GGTTAAAAAACGTCTAGGCCCCCCGAACCACGGGGACGTGGTTTTCCTTT TaqI

GAAAAACACG<u>ATT</u>G<u>C</u>-3'
XhoI (SEQ ID NO: 7)

5

10

20

30

35

40

50

55

This sequence matches the EMC virus leader sequence from nucleotide 763 to 827. It also changes the ATG at position 10 within the EMC virus leader to an ATT and is followed by a Xhol site. A three way ligation of the pMT21 Eco RI-Xhol fragment, the EMC virus EcoRI-Taql fragment, and the 68 bp oligonucleotide adapter Taql-Xhol adapter resulting in the vector pEMC2\(\beta\)1.

This vector contains the SV40 origin of replication and enhancer, the adenovirus major late promoter, a cDNA copy of the majority of the adenovirus tripartite leader sequence, a small hybrid intervening sequence, an SV40 polyadenylation signal and the adenovirus VA I gene, DHFR and β-lactamase markers and an EMC sequence, in appropriate relationships to direct the high level expression of the desired cDNA in mammalian cells.

The construction of vectors may involve modification of the BMP-9 DNA sequences. For instance, BMP-9 cDNA can be modified by removing the non-coding nucleotides on the 5' and 3' ends of the coding region. The deleted non-coding nucleotides may or may not be replaced by other sequences known to be beneficial for expression. These vectors are transformed into appropriate host cells for expression of BMP-9 proteins.

One skilled in the art can manipulate the sequences of Figure 1 or Figure 3 (SEQ ID NO: 1 and 8) by eliminating or replacing the mammalian regulatory sequences flanking the coding sequence with bacterial sequences to create bacterial vectors for intracellular or extracellular expression by bacterial cells. For example, the coding sequences could be further manipulated (e.g. ligated to other known linkers or modified by deleting non-coding sequences therefrom or altering nucleotides therein by other known techniques). The modified BMP-9 coding sequence could then be inserted into a known bacterial vector using procedures such as described in T. Taniguchi et al., <u>Proc. Natl Acad. Sci. USA</u>, 77:5230-5233 (1980). This exemplary bacterial vector could then be transformed into bacterial host cells and a BMP-9 protein expressed thereby. For a strategy for producing extracellular expression of BMP-9 proteins in bacterial cells, see, e.g. European patent application EPA 177,343.

Similar manipulations can be performed for the construction of an insect vector [See, e.g. procedures described in published European patent application 155,476] for expression in insect cells. A yeast vector could also be constructed employing yeast regulatory sequences for intracellular or extracellular expression of the factors of the present invention by yeast cells. [See, e.g., procedures described in published PCT application W086/00639 and European patent application EPA 123,289].

A method for producing high levels of a BMP-9 protein of the invention in mammalian cells may involve the construction of cells containing multiple copies of the heterologous BMP-9 gene. The heterologous gene is linked to an amplifiable marker, e.g. the dihydrofolate reductase (DHFR) gene for which cells containing increased gene copies can be selected for propagation in increasing concentrations of methotrexate (MTX) according to the procedures of Kaufman and Sharp, <u>J. Mol. Biol.</u>, 159:601-629 (1982). This approach can be employed with a number of different cell types.

For example, a plasmid containing a DNA sequence for a BMP-9 of the invention in operative association with other plasmid sequences enabling expression thereof and the DHFR expression plasmid pAdA26SV(A)3 [Kaufman and Sharp, Mol. Cell. biol., 2:1304 (1982)] can be co-introduced into DHFR-deficient CHO cells, DUKX-BII, by various methods including calcium phosphate coprecipitation and transfection, electroporation or protoplast fusion. DHFR expressing transformants are selected for growth in alpha media with dialyzed fetal calf serum, and subsequently selected for amplification by growth in increasing concentrations of MTX (e.g. sequential steps in 0.02, 0.2, 1.0 and 5uM MTX) as described in Kaufman et al., Mol Cell Biol., 5:1750 (1983). Transformants are cloned, and biologically active BMP-9 expression is monitored by the Rosen-modified Sampath - Reddi rat bone formation assay described above in Example III. BMP-9 expression should increase with increasing levels of MTX resistance. BMP-9 polypeptides are characterized using standard techniques known in the art such as pulse labeling with [35S] methionine or cysteine and polyacrylamide gel electrophoresis. Similar procedures can be followed to produce other related BMP-9 proteins.

A. BMP-9 Vector Construction

In order to produce human BMP-9 proteins of the invention DNA sequences encoding the mature region of the human BMP-9 protein may be joined to DNA sequences encoding the propertide region of the murine BMP-9 protein. This murine/human hybrid DNA sequence is inserted into an appropriate expression vector and introduced into mammalian cells or other preferred eukaryotic or prokaryotic hosts by conventional genetic engineering techniques. The

construction of this murine/human BMP-9 containing expression plasmid is described below.

A derivative of the human BMP-9 sequence (SEQ ID NO:8) comprising the nucleotide sequence from nucleotide #105 to #470 is specifically amplified. The following oligonucleotides are utilized as primers to allow the amplification of nucleotides #105 to #470 of the human BMP-9 sequence (SEQ ID NO:8) from clone pGEM-111 described above.

5

#3 ATCGGGCCCCTTTTAGCCAGGCGGAAAAGGAG

#4 AGCGAATTCCCCGCAGGCAGATACTACCTG

10

This procedure generates the insertion of the nucleotide sequence ATCGGGCCCCT immediately preceeding nucleotide #105 and the insertion of the nucleotide sequence GAATTCGCT immediately following nucleotide #470. The addition of these sequences results in the creation of an Apa I and EcoR I restriction endonuclease site at the respective ends of the specifically amplified DNA fragment. The resulting 374 bp Apa I/EcoR I fragment is subcloned into the plasmid vector pGEM-7Zf(+) (Promega catalog# p2251) which has been digested with Apa I and EcoR I. The resulting clone is designated phBMP9mex-1.

The following oligonucleotides are designed on the basis of murine BMP-9 sequences (SEQ ID NO:1) and are modified to facilitate the construction of the murine/human expression plasmid referred to above:

20

#5

GATTCCGTCGACCACCATGTCCCCTGGGGCCTGGTCTAGATGGATACACAGCTGTGGGGCC

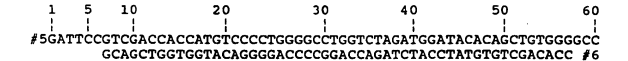
25

30

#6 CCACAGCTGTGTATCCATCTAGACCAGGCCCCAGGGGACATGGTGGTCGACG

These oligonucleotides contain complimentary sequences which upon addition to each other facilitate the annealing (base pairing) of the two individual sequences, resulting in the formation of a double stranded synthetic DNA linker (designated LINK-1) in a manner indicated below:

35



40

50

55

This DNA linker (LINK-1) contains recognition sequences of restriction endonucleases needed to facilitate subsequent manipulations required to construct the murine/human expression plasmid, as well as sequences required for maximal expression of heterologous sequences in mammalian cell expression systems. More specifically (referring to the sequence numbering of oligonucleotide #5/LINK-1); nucleotides #1-#11 comprise recognition sequences for the restriction endonucleases BamH I and Sal I, nucleotides #11-#15 allow for maximal expression of heterologuos sequences in mammallian cell expression systems, nucleotides #16-#31 correspond to nucleotides #610-#625 of the murine BMP-9 sequence (SEQ ID NO:1), nucleotides #32-#33 are inserted to facilitate efficient restriction digestion of two adjacent restriction endonuclease sites (EcoO109 I and Xba I), nucleotides #34-#60 correspond to nucleotides #1515-#1541 of the murine BMP-9 sequence (SEQ ID NO:1) except that nucleotide #58 of synthetic oligonucloetide #5 is a G rather than the A which appears at position #1539 of SEQ ID NO:1 (This nucleotide conversion results in the creation of an Apa I restriction endonuclease recognition sequence, without altering the amino acid sequence it is intended to encode, to facilitate further manipulations of the murine/human hybrid expression plasmid. LINK-1 (the double stranded product of the annealing of oligonucleotides #5 and #6) is subcloned into the plasmid vector pGEM-7Zf(+) which has been digested with the restriction endonucleases Apa I and BamH I. This results in a plasmid in which the sequences normally present between the Apa I and BamH I sites of the pGEM-7Zf(+) plasmid polylinker are replaced with the sequences of LINK-1 described above. The resulting plasmid clone is designated pBMP-9link.

pBMP-9link is digested with the restriction endonucleases BamH I and Xba I resulting in the removal nucleotides #1-#34 of LINK-1 (refer to the numbering of oligo #5). Clone ML14a, which contains an insert comprising the sequence

set forth in SEQ ID NO:1, is also digested with the restriction endonucleases BamH I and Xba I resulting in the removal of sequences comprising nucloetides #1-#1515 of SEQUENCE ID NO:1 (murine BMP-9). This BamH I/Xba I fragment of mouse BMP-9 is isolated from the remainder of the ML14a plasmid clone and subcloned into the BamH I/Xba I sites generated by the removal of the synthetic linker sequences described above. The resulting clone is designated p302.

5

20

25

30

35

40

45

50

The p302 clone is digested with the restriction endonuclease EcoO109 I resulting in the excision of nucloetides corresponding to nucleotides #621-#1515 of the murine BMP-9 sequence (SEQ ID NO:1) and nucleotides #35-#59 of LINK-1 (refer to numbering of oligonucleotide #5). It should be noted that the Apa I restriction site created in LINK-1 by the A to G conversion described above is a subset of the recognition sequence of EcoO109 I, therefore digestion of p302 with EcoO109 I cleaves at the Apa I site as well as the naturally occuring murine EcoO109 I (location #619-#625 of SEQ ID NO:1) resulting in the excision of a 920 bp EcoO109 I/EcoO109 I (Apa I) fragment comprising the sequences described above. This 920 EcoO109 I/EcoO109 I (Apa I) fragment is isolated from the remainder of the p302 plasmid clone and subcloned into clone pBMP-9link which has been similarly digested with EcoO109 I. It should be noted that the nucleotides GG (#32-#33 of oligonucleotide #5) originally designed to facilitate a more complete digestion of the two adjacent restriction sites EcoO109 I and Xba I of LINK-1, which is now a part of pBMP-9link (described above), results in the creation of Dcm methylation recognition sequence. The restriction nuclease EcoO109 I is sensitive to Dcm methylation and therefore cleavage of this sequence (nucleotides #25-#31 of oligonucleotide #5/LINK-1) by the restriction endonuclease EcoO109 I is prevented at this site. Therefore the plasmid clone pBMP-9link is cleaved at the Apa I site but not at the EcoO109 I site upon digestion with the restriction endonuclease EcoO109 I as described above, preventing the intended removal of the sequences between the EcoO109 I and Xba I site of LINK-1 (#32-#55 defined by the numbering of oligonucleotide #5). This results in the insertion of the 920 bp EcoO109 I/Apa I fragment at the EcoO109 I (Apa I) site of pBMP-9link. The resulting clone is designated p318.

Clone p318 is digested with the restriction endonucleases Sal I and Apa I, resulting in the excision of sequences comprising nucleotides #6-#56 of LINK-1 (refer to oligo #5 for location), nucleotides #621-#1515 of murine BMP-9 (SEQ ID NO:1), and nucleotides #35-#60 of LINK-1 (refer to oligo #5 for location). The resulting 972 bp Sal I/Apa I fragment described above is isolated from the remainder of the p318 plasmid clone and will be utilized in subsequent manipulations.

The clone phBMP9mex-1 (described above), which contains DNA sequences which encode the entire mature region and portions of the propeptide of the human BMP-9 protein, is digested with the restriction endonucleases Apa I and EcoR I. This results in the excision of a 374 bp fragment comprising nucleotides #105-#470 of the human BMP-9 sequence (SEQ ID NO:8) and the additional nucleotides of oligonucleotide primers #3 and #4 which contain the recognition sequences for the restriction endonucleases Apa I and EcoR I. This 374 bp Apa I/EcoR I fragment is combined with the 972 bp Sal I/Apa I fragment from p138 (isolation described above) and ligated to the mammalian cell expression plasmid pED6 (a derivative of pEMC2β1) which has been digested with Sal I and EcoR I. The resulting clone is designated p324.

The clone ML14a (murine BMP-9) is digested with EcoO109 I and Xba I to generate a fragment comprising nucleotides #621-#1515 of SEQ ID NO:1.

The following oligonucleotides are synthesized on an automated DNA synthesizer and combined such that their complimentary sequences can base pair (anneal) with each other to generate a double stranded synthetic DNA linker designated LINK-2:

#7 TCGACCACCATGTCCCCTGG

#8 GCCCCAGGGGACATGGTGG

This double stranded synthetic DNA linker (LINK-2) anneals in such a way that it generates single stranded ends which are compatible to DNA fragments digested with Sal I (one end) or EcoO109 I (the other end) as indicated below:

#7 TCGACCACCATGTCCCCTGG GGTGGTACAGGGGACCCCG #8:

This LINK-2 synthetic DNA linker is ligated to the 895 bp EcoO109 I/Xba I fragment comprising nucleotides #621-#1515 of murine BMP-9 (SEQ ID NO:1) described above. This results in a 915 bp Sal I/Xba I fragment.

The clone p324 is digested with Sal I/Xba I to remove sequences comprising nucleotides #6-#56 of LINK-1 (refer to oligo #5 for location) and nucleotides #621-#1515 of murine BMP-9 (SEQ ID NO:1). The sequences comprising

nucleotides #35-#60 of LINK-1 (refer to oligo #5 for location) and the sequences comprising the 374 bp Apa I/EcoR I fragment (human BMP-9 sequences) derived from phBMP9mex-1 remain attached to the pED6 backbone. The 915 bp Sal I/Xba I fragment comprising LINK-2 sequences and nucleotides #621-#1515 of murine BMP-9 (SEQ ID NO:1) is ligated into the p324 clone from which the Sal I to Xba I sequences described above have been removed.

The resulting plasmid is designated BMP9fusion and comprises LINK-2, nucleotides #621-#1551 of murine BMP-9 (SEQ ID NO:1), nucleotides #35-#59 of LINK-1 (refer to the numbering of oligonucleotide #5), and the 374 bp Apa I/EcoR I fragment (human BMP-9) derived from clone pBMP9mex-1 (described above) inserted between the Sal I and EcoR I sites of the mammalian cell expression vector pED6.

BMP9 fusion is transfected into CHO cells using standard techniques known to those having ordinary skill in the art to create stable cell lines capable of expressing human BMP-9 protein. The cell lines are cultured under suitable culture conditions and the BMP-9 protein is isolated and purified from the culture medium.

EXAMPLE V

5

10

20

25

35

45

50

55

15 Biological Activity of Expressed BMP-9

To measure the biological activity of the expressed BMP-9 proteins obtained in Example IV above, the proteins are recovered from the cell culture and purified by isolating the BMP-9 proteins from other proteinaceous materials with which they are co-produced as well as from other contaminants. The purified protein may be assayed in accordance with the rat bone formation assay described in Example III.

Purification is carried out using standard techniques known to those skilled in the art. It is contemplated, as with other BMP proteins, that purification may include the use of Heparin sephanose.

Protein analysis is conducted using standard techniques such as SDS-PAGE acrylamide [U.K. Laemmli, Nature 227:680 (1970)] stained with silver [R.R. Oakley, et al. Anal. Biochem. 105:361 (1980)] and by immunoblot [H. Towbin, et al. Proc. Natl. Acad. Sci. USA 76:4350 (1979)]

The foregoing descriptions detail presently preferred embodiments of the present invention. Numerous modifications and variations in practice thereof are expected to occur to those skilled in the art upon consideration of these descriptions.

30 SEQUENCE LISTING

- (1) GENERAL INFORMATION:
 - (i) APPLICANT: Wozney, John M. Celeste, Anthony
 - (ii) TITLE OF INVENTION: BMP-9 COMPOSITIONS
 - (iii) NUMBER OF SEQUENCES: 9
- 40 (iv) CORRESPONDENCE ADDRESS:
 - (A) ADDRESSEE: Genetics Institute, Inc.
 - (B) STREET: Legal Affairs 87 CambridgePark Drive
 - (C) CITY: Cambridge
 - (D) STATE: MA
 - (E) COUNTRY: US
 - (F) ZIP: 02140
 - (v) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk
 - (B) COMPUTER: IBM PC compatible
 - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
 - (D) SOFTWARE: Patentin Release #1.0, Version #1.25
 - (vi) CURRENT APPLICATION DATA:
 - (A) APPLICATION NUMBER: US

	(B) FILING DATE: (C) CLASSIFICATION:	
5	(viii) ATTORNEY/AGENT INFORMATION:	
5	(A) NAME: Kapinos, Ellen J.(B) REGISTRATION NUMBER: 32,245(C) REFERENCE/DOCKET NUMBER: GI 5186A	
10	(ix) TELECOMMUNICATION INFORMATION:	
	(A) TELEPHONE: (617) 876-1170 (B) TELEFAX: (617) 876-5851	
15	(2) INFORMATION FOR SEQ ID NO:1:	
	(i) SEQUENCE CHARACTERISTICS:	
20	(A) LENGTH: 2447 base pairs(B) TYPE: nucleic acid(C) STRANDEDNESS: double(D) TOPOLOGY: linear	
05	(ii) MOLECULE TYPE: cDNA to mRNA	
25	(iii) HYPOTHETICAL: NO	
	(iv) ANTI-SENSE: NO	
30	(vi) ORIGINAL SOURCE:	
35	(A) ORGANISM: Mus musculus (B) STRAIN: C57B46xCBA (F) TISSUE TYPE: liver	
00	(vii) IMMEDIATE SOURCE:	
40	(A) LIBRARY: Mouse liver cDNA (B) CLONE: ML14A	
	(viii) POSITION IN GENOME:	
	(C) UNITS: bp	
45	(ix) FEATURE:	
	(A) NAME/KEY: mat_peptide (B) LOCATION: 15641893	
50	(ix) FEATURE:	
	(A) NAME/KEY: CDS (B) LOCATION: 6101896	
55	(ix) FEATURE:	
	(A) NAME/KEY: mRNA (B) LOCATION: 12447	

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

5	CATTAATAAA TATTAAGTAT TGGAATTAGT GAAATTGGAG TTCCTTGTGG AAGGAAGTGG	60
	GCAAGTGAGC TTTTTAGTTT GTGTCGGAAG CCTGTAATTA CGGCTCCAGC TCATAGTGGA	120
	ATGGCTATAC TTAGATTTAT GGATAGTTGG GTAGTAGGTG TAAATGTATG TGGTAAAAGG	180
10	CCTAGGAGAT TTGTTGATCC AATAAATATG ATTAGGGAAA CAATTATTAG GGTTCATGTT	240
	CGTCCTTTTG GTGTGTGGAT TAGCATTATT TGTTTGATAA TAAGTTTAAC TAGTCAGTGT	300
	TGGAAAGAAT GGAGACGGTT GTTGATTAGG CGTTTTGAGG ATGGGAATAG GATTGAAGGA	360
15	AATATAATGA TGGCTACAAC GATTGGGAAT CCTATTATTG TTGGGGTAAT GAATGAGGCA	420
	AATAGATTTT CGTTCATTTT AATTCTCAAG GGGTTTTTAC TTTTATGTTT GTTAGTGATA	480
	TTGGTGAGTA GGCCAAGGGT TAATAGTGTA ATTGAATTAT AGTGAAATCA TATTACTAGA	540
20	CCTGATGTTA GAAGGAGGC TGAAAAGGCT CCTTCCCTCC CAGGACAAAA CCGGAGCAGG	600
	GCCACCCGG ATG TCC CCT GGG GCC TTC CGG GTG GCC CTG CTC CCG CTG Met Ser Pro Gly Ala Phe Arg Val Ala Leu Leu Pro Leu -318 -315 -310	648
25	TTC CTG CTG GTC TGT GTC ACA CAG CAG AAG CCG CTG CAG AAC TGG GAA Phe Leu Leu Val Cys Val Thr Gln Gln Lys Pro Leu Gln Asn Trp Glu -305 -290	696
30	CAA GCA TCC CCT GGG GAA AAT GCC CAC AGC TCC CTG GGA TTG TCT GGA Gln Ala Ser Pro Gly Glu Asn Ala His Ser Ser Leu Gly Leu Ser Gly -285 -280 -275	744
	GCT GGA GAG GAG GGT GTC TTT GAC CTG CAG ATG TTC CTG GAG AAC ATG Ala Gly Glu Gly Val Phe Asp Leu Gln Met Phe Leu Glu Asn Met -270 -265 -260	792
35	AAG GTG GAT TTC CTA CGC AGC CTT AAC CTC AGC GGC ATT CCC TCC CAG Lys Val Asp Phe Leu Arg Ser Leu Asn Leu Ser Gly Ile Pro Ser Gln	840

(2) INFORMATION FOR SEQ ID NO:9:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 151 amino acids

(B) TYPE: amino acid (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:

55

40

45

	* -41	Thr -40	Arg	Glu	Сув	Ser	Arg -35	Ser	Сув	Pro	Arg	Thr	Ala	Pro	Gln	Arg
5	Gln -25	Val	Arg	Ala	Val	Thr	Arg	Arg	Thr	Arg	Met -15	Ala	His	Val	Ala	Ala -10
	Gly	Ser	Thr	Leu	Ala -5	Arg	Arg	Lys	Arg	Ser 1	Ala	Gly	Ala	Gly 5	Ser	His
10	Сув	Gln	Lys 10	Thr	Ser	Leu	Arg	Val 15	Asn	Phe	Glu	Asp	Ile 20	Gly	dıı	Хвр
	Ser	Trp 25	Ile	Ile	Ala	Pro	Lys 30	Glu	Tyr	Glu	Ala	Tyr 35	Glu	Сув	Lys	Gly
15	Gly 40		Phe	Phe	Pro	Leu 45	Ala	Asp	Asp	Val	Thr 50	Pro	Thr	Lys	His	Ala 55
	Ile	Val	Gln	Thr	Leu 60	Val	His	Leu	Lys	Phe 65	Pro	Thr	Lys	Val	Gly 70	Lys
20	Ala	Сув	Cys	Val 75		Thr	Lys	Leu	Ser 80		Ile	Ser	Val	Leu 85	Tyr	Lys
	qaA	Asp	Met 90		Val	Pro	Thr	Leu 95		Tyr	His	Tyr	Glu 100		Met	Ser
25	Val	Ala 105		Cys	Gly	Сув	Arg 110									
30	(ix)	FEAT	URE:													
		, ,		KEY: e												
35	(ix)	FEAT														
				KEY: C												
40	(ix)	FEAT														
				ΚΕΥ: π ΙΟΝ: 1	-	-										
45	(ix)	FEAT														
				KEY: m												
50	(xi)	` ,		E DES		ION:	SEQ II	D NO:	8:							

	*	ACA Thr -40			TGC Cys												48
5	CAG Gln -25	GTG Val	AGA Arg	GCA Ala	GTC Val	ACG Thr -20	AGG Arg	AGG Arg	ACA Thr	CGG Arg	ATG Met -15	GCG Ala	CAC His	GTG Val	GCT Ala	GCG Ala -10	96
10	GGG Gly	TCG Ser	ACT Thr	TTA Leu	GCC Ala -5	AGG Arg	CGG Arg	ГÀе УУУ	AGG Arg	AGC Ser 1	GCC Ala	GGG Gly	GCT Ala	GGC Gly 5	AGC Ser	CAC His	144
		CAA Gln		Thr													192
15	AGC Ser	TGG Trp 25	Ile	ATT Ile	GCA Ala	CCC	AAG Lys 30	Glu	TAT Tyr	GAA Glu	GCC Ala	TAC Tyr 35	GAG Glu	TGT Cys	AAG Lys	GGC Gly	240
20	GG(Gl _y 4(TGC Cys	TTC Phe	TTC Phe	CCC Pro	TTG Leu 45	Ala	GAC	GAT Asp	GTG Val	ACG Thr 50	Pro	ACG Thr	AAA Lys	CAC His	GCT Ala 55	288
	ATC Ile	GTG Val	CAG Gln	ACC Thr	CTG Leu 60	Val	CAT His	CTC Leu	AAG Lys	TTC Phe 65	Pro	ACA Thr	AAG Lys	GTG Val	GGC Gly 70		336
25	GC: Ala	TGC Cys	TGT Cys	GTG Val	Pro	ACC	AAA Lys	CTG Lev	AGC Ser 80	Pro	ATC	TCC Ser	GTC Val	CTC Lev 85	Tyr	AAG Lys	384
30	GA' As	T GAG p Asi	ATC Met	: Gly	GTG Val	Pro	ACC Thr	CTC Lev	Lys	TAC Tyr	CAT His	TAC Tyr	GAG Glu 100	ı Gly	ATG Met	AGC Ser	432
		G GCI 1 Ala 10	a Glu					3	TATO	TGC	CTG	CGGG					470
35								Cam	egg	CAGO	· mc	CAC					
								CAI		CAGC		GAG					
40	(2) INF	ORMA	TION	FOR	SEQ I	D NO	:6:										
	(i)	SEQU	ENCE	CHA	RACT	ERIS	TICS:										
45		(B) T (C) S	YPE:	nuclei IDEDI	base ic acid NESS linea	doub	ole										
50	(ii)	MOLE	CULE	TYP	E: cDI	NA to	mRN.	A									
50	(xi)	SEQI	JENC	E DES	SCRIP	TION	: SEC	N DI Q	O:6:								
55	CTGC	AGGCG	GA GC	CTG	ATTO	CT(CGAG	CCAT	CAT	G							34
	(2) INF	ORMA	TION	FOR	SEQI	D NO	:7:										

	(i) SEQUENCE CHARACTERISTICS:	
5	(A) LENGTH: 68 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: double	
	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: cDNA to mRNA	
10	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:	
	CGAGGTTAAA AAACGTCTAG GCCCCCCGAA CCACGGGGAC GTGGTTTTCC TTTGAAAAAC	60
15	ACGATTGC	68
	(2) INFORMATION FOR SEQ ID NO:8:	
20	(i) SEQUENCE CHARACTERISTICS:	
20	(A) LENGTH: 470 base pairs	
	(B) TYPE: nucleic acid (C) STRANDEDNESS: double	
	(D) TOPOLOGY: linear	
25	(ii) MOLECULE TYPE: DNA (genomic)	
	(iii) HYPOTHETICAL: NO	
30	(v) FRAGMENT TYPE: C-terminal	
	(vi) ORIGINAL SOURCE:	
35	(A) ORGANISM: Homo sapiens (H) CELL LINE: W138 (genomic DNA)	
	(vii) IMMEDIATE SOURCE:	
40	(A) LIBRARY: human genomic library (B) CLONE: lambda 111-1	
	(viii) POSITION IN GENOME:	
45	(C) UNITS: bp	
50		
55		

	Arg	Ile -115	naA	Ile	Tyr	Glu	Val -11	Met .0	Lys	Pro	Pro		G1u L05	Val	Val	Pro
5	Gly -100	His	Leu	Ile	Thr	Arg -95	Leu	Leu	Asp	Thr	Arg -9(Val	His	His	Asn -85
	Val	Thr	Arg	Trp	Glu -80	Thr	Phe	Asp	Val	Ser -75	Pro	Ala	Val	Leu	Arg -70	Trp
10	Thr	Arg	Glu	Lys -65	Gln	Pro	Asn	Tyr	Gly -60	Leu	Ala	Ile	Glu	Val -55	Thr	His
15	Leu	His	Gln -50	Thr	Arg	Thr	His	Gln -45	Gly	Gln	His	Vál	Arg -40	Ile	Ser	Arg
	Ser	Leu -35	Pro	Gln	Gly	Ser	Gly -30	naA	Trp	Ala	Gln	Leu -25	Arg	Pro	Leu	Leu
20	Val -20	Thr	Phe	Gly	His	Asp -15	Gly	Arg	Gly	His	Ala -10	Leu	Thr	Arg	Arg	Arg -5
	Arg	Ala	Lys	Arg	Ser 1	Pro	Lys	His	His 5	Ser	Gln	Arg	Ala	Arg 10	Lys	Lys
25	Asn	Lys	Asn 15	Cys	Arg	Arg	His	Ser 20	Leu	Tyr	Val	qaƙ	Phe 25	Ser	qaƙ	Val
	Gly	Trp 30	Asn	Asp	Trp	Ile	Val 35	Ala	Pro	Pro	Gly	Tyr 40	Gln	Ala	Phe	Tyr
30 e ama	Cys 45	His	Gly	Asp	Сув	Pro 50	Phe	Pro	Leu	Ala	Asp 55	His	Leu	Asn	Ser	Thr 60
	Asn	His	Ala	Ile	Val 65	Gln	Thr	Leu	Val	Asn 70	Ser	Val	Asn	Ser	Ser 75	Ile
35	Pro	Lys	Ala	ayo 80	Сув	Val	Pro	Thr	Glu 85	Leu	Ser	Ala	Ile	Ser 90	Met	Leu
40	Tyr	Leu	Asp 95	Glu	Tyr	Asp	Lys	Val 100	Val	Leu	Lys	Asn	Tyr 105	Gln	Glu	Met
, ,	Val	Val 110	Glu	Gly	Суѕ	Gly	Cys 115	Arg								

(2) INFORMATION FOR SEQ ID NO:5:

45

50

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 15 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: cDNA to mRNA
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:

	TGT GGG TGC C Cys Gly Cys A 115		CAGTCCTTGA GGAT.	AGACAG ATATACACAC	1666
5	CACACACACA CA	ACCACATAC ACCA	CACACA CACGTTCCC	A TCCACTCACC CACAC	ACTAC 1726
	ACAGACTGCT TO	CTTATAGC TGGA	CTTTTA TTTAAAAA	A AAAAAAAAA AATGG	AAAAA 1786
	ATCCCTAAAC AT	TCACCTTG ACCI	TATTA TGACTTAC	G TGCAAATGTT TTGAC	CATAT 1846
10	TGATCATATA TT	rtigacaaa atai	PATTTAT AACTACGTA	T TAAAAGAAAA AAATA	AAATG 1906
	AGTCATTATT TT	KAAA AAAAAAA	AAAAACT CTAGAGTCG	A CGGAATTC	1954
15	(2) INFORMATIO	N FOR SEQ ID NO:4	4 :		
	(i) SEQUENC	E CHARACTERIST	TICS:		
20	(B) TYPE	GTH: 408 amino acid E: amino acid DLOGY: linear	ls		
	(ii) MOLECUL	LE TYPE: protein			
25	(xi) SEQUEN	CE DESCRIPTION:	SEQ ID NO:4:		
					•
	Met Ile Pro (-292 -29		et Leu Met Val Va -285	al Leu Leu Cys Gln -280	Val
30	Leu Leu Gly -275		is Ala Ser Leu II -270	le Pro Glu Thr Gly -265	Lys
	Lys Lys Val -260	Ala Glu Ile G -255		ly Gly Arg Arg Ser -250	Gly -245
35	Gln Ser His	Glu Leu Leu A -240	rg Asp Phe Glu A -235	la Thr Leu Leu Gln -2	Met 30
	Phe Gly Leu	Arg Arg Arg P -225	Pro Gln Pro Ser Ly -220	ys Ser Ala Val Ile -215	Pro
40	Asp Tyr Met -210		Tyr Arg Leu Gln So -205	er Gly Glu Glu Glu -200	Glu
	Glu Gln Ile -195	His Ser Thr G	Gly Leu Glu Tyr P -190	ro Glu Arg Pro Ala ~185	Ser
45	Arg Ala Asn -180	Thr Val Arg S		lu Glu His Leu Glu -170	Asn -165
	Ile Pro Gly	Thr Ser Glu A	Asn Ser Ala Phe A -155	rg Phe Leu Phe Asn	Leu 150
50	Ser Ser Ile	Pro Glu Asn C	Glu Val Ile Ser S -140	er Ala Glu Leu Arg -135	Leu
55	Phe Arg Glu -130		Gln Gly Pro Asp T -125	rp Glu Arg Gly Phe -120	His

	GTG Val	GAC Asp	CAG Gln	GGC Gly -125	CCT Pro	GAT Asp	TGG Trp	GAA Glu	AGG Arg -120	Gly	TTC Phe	CAC His	CGT Arg	ATA Ile -115	Asn	ATT Ile	942
5				Met	AAG Lys				Glu					aiH			990
10					GAC Asp												1038
	GAA Glu -80	ACT Thr	TTT Phe	GAT Asp	GTG Val	AGC Ser -75	CCT Pro	GCG Ala	GTC Val	CTT Leu	CGC Arg -70	TGG Trp	ACC Thr	CGG Arg	GAG Glu	AAG Lys -65	1086
15					GGG Gly -60												1134
20					GGC Gly												1182
					TGG Trp												1230
25					GGC Gly												1278
30					CAC His 5												1326
					CTC Leu												1374
35					CCA Pro				Gln								1422
40			Phe		CTG Leu			His					Asn				1470
		Gln			GTC Val		Ser					Ile					1518
45						Leu					Met					GAG Glu	1566
50					. Val					Gln					Glu	GGA Gly	1614

(B) LOCATION: 9..1934

55

(xi) SAQUENCE DESCRIPTION: SEQ ID NO:3:

	CTCTAGA	GGG C	AGAGO	BAGGA	GGG	AGGC	BAGG	GAA	GGAG	CGC	GGAG	CCCG	GC C	CGGA	AGCTA	60
	GGTGAGT	GTG G	CATC	CGAGC	TGA	GGG7	ACGC	GAG	CCTG	AGA	CGCC	GCTG	CT G	CTCC	GGCTG	120
5	AGTATCT	AGC I	TGTCT	recec	GAT	GGG2	ATTC	CCG!	TCCA	AGC	TATC	TCGA	GC C	TGCA	GCGCC	180
	ACAGTCC	CCG G	CCCT	CGCCC	AGG	TTC	ACTG	CAA	CCGT	TCA	GAGG	TCCC	CA G	GAGC	TGCTG	240
	CTGGCGA	GCC C	GCTA	CTGCA	GGG	ACC!	PTATG	GAG	CCAT	TCC	GTAG	TGCC	AT C	CCGA	GCAAC	300
10	GCACTGC	TGC A	GCTT	cccr	AGC	CTT:	rcca	GCA	AGTT	TGT	TCAA	GATT	GG C	TGTC	AAGAA	360
	TCATGGA	CTG 1	TATT	ATATO	CCT	TGT.	PTTC	TGT	CAAG	ACA	M		le P			414
15	AAC CGA	ATG Met	CTG : Leu : -285	Met \	STC G /al V	TT !	Leu	TTA Leu -280	Сув	CAA Gln	GTC Val	CTG Leu	CTA Leu -275	Gly	GGC Gly	462
20	GCG AGO Ala Ser	CAT His	Ala	AGT 3 Ser 1	MG A Leu I	le:	CCT Pro -265	Glu	ACG Thr	GGG Gly	AAG Lys	AAA Lys -260	Lys	GTC Val	GCC Ala	510
٠.	GAG ATT	e Gln			Ala (Gly					Gln				558
25	CTC CTC Leu Leu -240			Phe (Met					606
30	Arg Ar	C CCG g Pro	CAG Gln	CCT Pro	Ser 1	AAG Lys	AGT Ser	GCC Ala	GTC Val -215	Ile	CCG Pro	GAC Asp	TAC Tyr	ATG Met -210	Arg	654
	GAT CT			Leu					Glu					Ile		. 702
35	AGC AC		Leu					Arg					Ala			750
40	GTG AG Val Ar	g Ser			His		Glu					Ile				798
	AGT GA Ser Gl -160					Arg					Leu					846
45	GAG AA Glu As			Ile		Ser	Ala	Glu	Leu	Arg					Gln	894

					-10					- 5					1	
5	Gly	Ala	Ser 5	Ser	His	Cys	Gln	Lys 10	Thr	Ser	Leu	Arg	Val 15	Asn	Phe	Glu
	Asp	Ile 20	Gly	Trp	Asp	Ser	Trp 25	Ile	Ile	Ala	Pro	Lys 30	Glu	Tyr	Asp	Ala
10	Tyr 35	Glu	Сув	Lys	Gly	Gly 40	Сув	Phe	Phe	Pro	Leu 45	Ala	Asp	Asp	Val	Th: 50
	Pro	Thr	Lув	His	Ala 55	Ile	Val	Gln	Thr	Leu 60	Val	His	Leu	Glu	Phe 65	Pro
15	Thr	Lys	Val	Gly 70	Lys	Ala	Cys	Сув	Val 75	Pro	Thr	Lys	Leu	Ser 80	Pro	Ile
	Ser	Ile	Leu 85	Tyr	Lys	qaƙ	Asp	Met 90	Gly	Val	Pro	Thr	Leu 95	Lys	Tyr	His
20	Tyr	Glu 100	Gly	Met	Ser	Val	Ala	Glu	Cys	Gly	Cys	Arg				

- (2) INFORMATION FOR SEQ ID NO:3:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 1954 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: cDNA to mRNA
- 35 (iii) HYPOTHETICAL: NO

25

30

40

45

50

55

- (iv) ANTI-SENSE: NO
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo sapiens
 - (G) CELL TYPE: Osteosarcoma Cell Line
 - (H) CELL LINE: U-20S
- (vii) IMMEDIATE SOURCE:
 - (A) LIBRARY: U20S cDNA in Lambda gtlO
 - (B) CLONE: Lambda U20S-3
- (viii) POSITION IN GENOME:
 - (C) UNITS: bp
 - (ix) FEATURE:

(A) NAME/KEY: CDS (B) LOCATION: 403..1629

	(ix) FEATURE:
_	(A) NAME/KEY: mat_peptide (B) LOCATION: 12791626
5	(ix) FEATURE:
10	(A) NAME/KEY: mRNA (xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:
15	
20	
25	
30	
35	
40	
45	
50	

	-318	ser 3	Pro	Gly -31	Ala 15	Phe	Arg	Val	Ala	Leu 310	Leu	Pro	Leu	Phe	Leu -305	Leu	
5	Val	Cys	Val -300	Thr	Gln	Gln	Lys	Pro	Leu 95	Gln	Asn	Trp	Glu -	Gln 290	Ala	Ser	
	Pro	Gly -285	Glu 5	Asn	Ala	His	Ser -28	ser 30	Leu	Gly	Leu	Ser	Gly 275	Ala	Gly	Glu	
10	Glu -270	Gly)	Val	Phe	Asp	Leu -26	Gln 55	Met	Phe	Leu	Glu -	Asn 260	Met	Lys	Val		255
	Phe	Leu	Arg	Ser	Leu -250) Nan	Leu	Ser	Gly	Ile -2	Pro 45	Ser	Gln	qaA		Thr 240	
15	Arg	Ala	Glu	Pro -235	Pro	Gln	Tyr	Met	11e -23	Asp 0	Leu	Tyr	Asn		Tyr 225	Thr	
	Thr	Asp	Lys -220	Ser	Ser	Thr	Pro	Ala -21	Ser 5	Asn	Ile	Val	Arg	Ser	Phe	Ser	
20	Val	Glu -205	Asp 5	Ala	Ile	Ser	Thr -20	Ala 0	Ala	Thr	Glu		Phe 195	Pro	Phe	Gln	
	Lys -190	His)	Ile	Leu	Ile	Phe	Asn 35	Ile	Ser	Ile		Arg 180	His	Glu	Gln		175
25	Thr	Arg	Ala	Glu	Leu -170	Arg	Leu	Tyr	Val	Ser	Cys 55	Gln	Asn	Asp		Asp 60	
	Ser	Thr	His	Gly -155	Leu	Glu	Gly	Ser	Met -15	Val	Val	Tyr	Asp	Val	Leu 145	Glu	
30				Thr	•				-15	0			Thr	-]	145		
30	Asp	Ser	Glu -140 Asp	Thr	Trp	Asp	Gln	Ala -13 Gly	-15 Thr 5	Gly	Thr	Lys Leu	Thr -1	-) Phe .30	Leu	Val	
<i>30</i>	Asp	Ser Gln -125 Val	Glu -140 Asp	Thr	Trp Arg	Asp	Glu Glu -12 Arg	Ala -13 Gly	-15 Thr 5 Trp	Gly Glu	Thr Thr	Lys Leu -1	Thr -1 Glu	Phe .30 Val	Leu Ser	Val Ser Lys	95
35	Asp Ser Ala	Ser Gln -125 Val	Glu -14(Asp 5	Thr	Trp Arg	Asp Val	Glu -12 Arg	Ala -13 Gly 0	Thr 5 Trp Asp	Gly Glu Ser	Thr Thr Thr	Lys Leu -1 Thr	Thr -J Glu 115 Asn	Phe 30 Val	Leu Ser Asn	Val Ser Lys	95
	Asp Ser Ala -110	Gln -125 Val	Glu -140 Asp 5 Lys Val	Thr) Ile Arg	Trp Arg Trp Val -90	Asp Val -10 Gln	Glu -12 Arg)5 Ser	Ala -13 Gly 0 Ala	Thr 5 Trp Asp	Gly Glu Ser Glu -85	Thr Thr Thr Ser	Lys Leu -1 Thr 100 Cys	Thr -] Glu 115 Asn Asp	Phe 30 Val Lys	Leu Ser Asn Leu -80	Val Ser Lys -:	95
<i>35</i>	Asp Ser Ala -110 Leu Ile	Gln -125 Val Glu Ser	Glu -14(Asp Lys Val	Thr Ile Arg Thr	Trp Arg Trp Val -90	Asp Val -10 Gln Gly	Glu -12 Arg 5 Ser	Ala Gly O Ala His	Thr 5 Trp Asp Arg	Glu Ser Glu -85 Leu	Thr Thr Thr Ser	Lys Leu Thr 100 Cys	Thr —1 Glu 115 Asn Asp	Phe .30 Val Lys Thr	Leu Ser Asn Leu -80 Val	Val Ser Lys -: Asp	95
35	Asp Ser Ala -110 Leu Ile Ser	Gln -125 Val Glu Ser	Glu -140 Asp Lys Val Val	Thr Ile Arg Thr Pro -75	Trp Arg Trp Val -90 Pro	Asp Val -10 Gln Gly Asn	Glu Glu -12 Arg 05 Ser Ser	Ala Gly O Ala His Lys	Thr Trp Asp Arg Asn -70 Lys	Glu Ser Glu -85 Leu Glu	Thr Thr Thr Ser Pro	Lys Leu Thr 100 Cys Phe	Thr —] Glu 115 Asn Asp Phe Leu —50	Phe .30 Val Lys Thr Val -65	Leu Ser Asn Leu -80 Val	Val Ser Lys -! Asp Phe Lys	95
<i>35</i>	Asp Ser Ala -110 Leu Ile Ser Glu	Glu Ser Asn Met -45	Glu -140 Asp Lys Val Val Asp -60	Thr Ile Arg Thr Pro -75	Trp Arg Trp Val -90 Pro Ser	Asp Val -10 Gln Gly Asn Glu	Gln Glu -12 Arg 5 Ser Gly Gln -40	Ala Gly O Ala His Lys Thr	Thr Trp Asp Arg Arg Lys	Glu Ser Glu -85 Leu Glu Met	Thr Thr Thr Ser Pro Thr	Lys Leu Thr 100 Cys Phe Arg Val -35	Thr —I Glu 115 Asn Asp Phe Leu —50 Lys	Phe .30 Val Lys Thr Val -65 Glu	Leu Ser Asn Leu -80 Val Leu Ala	Val Ser Lys -! Asp Phe Lys	95

			-15					-10					- 5					
5	AGG Arg	AGC Ser 1	ACC Thr	GGA Gly	GCC Ala	AGC Ser 5	AGC Ser	CAC His	TGC Cys	CAG Gln	AAG Lys 10	ACT Thr	TCT Ser	CTC Leu	AGG Arg	GTG Val 15	1608	3
	AAC Asn	TTT Phe	GAG Glu	GAC Asp	ATC Ile 20	GGC Gly	TGG Trp	GAC Asp	AGC Ser	TGG Trp 25	ATC Ile	ATT Ile	GCA Ala	CCC Pro	AAG Lys 30	GAA Glu	1656	5
10	TAT Tyr	GAC Asp	GCC Ala	TAT Tyr 35	GAG Glu	TGT Cys	AAA Lys	GGG Gly	GGT Gly 40	TGC Cys	TTC Phe	TTC Phe	CCA Pro	TTG Leu 45	GCT Ala	GAT Asp	1704	1
15	GAC Asp	GTG Val	ACA Thr 50	Pro	ACC Thr	aaa Lys	CAT His	GCC Ala 55	ATC Ile	GTG Val	CAG Gln	ACC Thr	CTG Leu 60	GTG Val	CAT His	CTC Leu	1752	2
	GAG Glu	TTC Phe 65	Pro	ACA Thr	AAG Lys	GTG Val	GGC Gly 70	AAA Lys	GCC Ala	TGC Cys	TGC Cys	GTT Val 75	CCC Pro	ACC Thr	AAA Lys	CTG Leu	1800	0
20	AGT Ser 80	Pro	ATC Ile	TCC Ser	ATC Ile	CTC Leu 85	Tyr	AAG Lys	GAT Asp	GAC Asp	ATG Met 90	Gly	GTG Val	CCA Pro	ACC Thr	CTC Leu 95	184	8
25	AAG Lys	TAC Tyr	CAC His	TAT	GAG Glu 100	Gly	ATG Met	AGT Ser	GTG Val	GCT Ala 105	Glu	TGT Cys	GGG Gly	TGT Cys	AGG Arg 110	TAGTCC	CTGC	19(
	AGC	CACC	CAG	GGTG	GGGA	TA C	AGGA	CATG	G AA	GAGG	TTCT	GGT	ACGG'	TCC	TGCA	TCCTCC	196	3
	TGC	GCAT	GGT	ATGC	CTAA	GT T	GATC	AGAA	A CC	ATCC	TTGA	GAA	GAAA	AGG	AGTT.	AGTTGC	202	3
30	CCT	TCTT	GTG	TCTG	GTGG	GT C	CCTC	TGCT	G AA	GTGA	CAAT	GAC	TGGG	GTA	TGCG	GGCCTG	208	3
	TGG	GCAG	AGC	AGGA	GACC	CT G	GAAG	GGTT	A GT	GGGT	AGAA	AGA	TGTC	AAA	AAGG	AAGCTG	214	3
	TGG	GTAG	ATG	ACCI	GCAC	TC C	AGTG	ATTA	G AA	GTCC	AGCC	TTA	CCTG	TGA	GAGA	GCTCCT	220	3
35	GGC	ATCI	AAG	AGAA	CTCT	GC I	TCCT	CATC	A TC	CCCA	.CCGA	CTT	GTTC	TTC	CTTG	GGAGTG	226	3
	TGI	CCTC	AGG	GAGA	ACAG	CA T	TGCT	GTTC	C TG	TGCC	TCAA	GCT	CCCA	GCT	GACT	CTCCTG	232	3
	TGG	CTCA	TAG	GACI	GAAT	eg e	GTGA	.GGAA	G AG	CCTG	ATGC	CCT	CTGG	CAA	TCAG	AGCCCG	238	3
40	AAC	GACI	TCA	AAAC	CATCI	GG A	CAAC	TCTC	TT A	GACI	GATG	CTC	CAAC	ATA	ATTT	TTAAAA	244	3
	AGA	\G															244	7

(2) INFORMATION FOR SEQ ID NO:2:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 428 amino acids

(B) TYPE: amino acid (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein

55

50

			-255					-250					-245	,			
5	GAC A	\AA Lys -240	ACC Thr	AGA Arg	GCG Ala	Glu	CCA Pro -235	Pro	CAG Gln	TAC Tyr	ATG Met	ATC Ile -230	qaƙ	TTG Leu	TAC Tyr	AAC Asn	888
	AGA 1 Arg 1 -225				qaA		Ser					Ser					936
10	AGC 1					Asp					Ala					Phe	984
15	CCC !				His					Asn					Arg		1032
	GAG (Gln		Thr					Arg					Сув			1080
20	GAT (qaA					Leu					Val				1128
25	GTT Val -145	Leu					Thr					Thr					1176
	TTC Phe					Asp					Gly					Glu	1224
30	GTA Val				Val					Arg					Thr		1272
35				Leu	GAG Glu											GAC Asp	1320
					AGT Ser								Leu				1368
40					AAT Asn		Arg					Lys					1416
45					ATG Met -45	Ile					Glu					Lys	1464
					Ala					Gly					Glu	GAG Glu	1512

Claims

50

55

1. A DNA sequence encoding a protein having the biological activity of a BMP-9 protein of inducing the formation of

1560

GGT CTA GAT GGA TAC ACA GCT GTG GGA CCA CTT TTA GCT AGA AGG AAG Gly Leu Asp Gly Tyr Thr Ala Val Gly Pro Leu Ala Arg Arg Lys

cartilage and/or bone which sequence is

5

30

35

40

45

- (a) the DNA sequence from nucleotides 124 to 453 of SEQ ID No. 8; or
- (b) the DNA sequence from nucleotides 145 to 453 of SEQ ID No. 8; or
- (c) a DNA sequence which differs from the DNA sequence of (a) or (b) due to the degeneracies of the genetic code:
- (d) an allelic variant of the sequence of (a) or (b); or
- (e) a DNA sequence hybridizing under stringent conditions to the sequences of (a) or (b).
- A recombinant DNA molecule containing a DNA sequence according to claim 1.
 - 3. The recombinant DNA molecule according to claim 2 wherein said DNA sequence is under the control of regulatory elements allowing its expression in a desired host cell.
- 4. A host cell containing the recombinant DNA molecule according to claim 2 or 3.
 - 5. The host cell according to claim 4 which is a bacterial cell, a yeast cell or a mammalian cell.
- 6. A method for the production of a protein having the biological activity of a BMP-9 protein comprising the cultivation of a host cell according to claim 4 or 5 under conditions appropriate for expression of said DNA sequence and recovering said protein from the culture.
 - 7. A protein encoded by the DNA sequence of claim 1.
- 25 8. A protein produced by the method of claim 6.
 - 9. A protein having the biological activity of a BMP-9 protein comprising one of the following amino acid sequences
 - (a) the amino acid sequence from amino acids No. 8 to 110 as set forth in Fig. 3 (SEQ ID No. 9); or
 - (b) the amino acid sequence from amino acids No. 1 to 110 as set forth in Fig. 3 (SEQ ID No. 9).
 - 10. A protein having the biological activity of a BMP-9 protein wherein said protein is a dimer wherein each subunit comprises at least the amino acid sequence from amino acids No. 8 to 110 of Fig. 3 (SEQ ID No. 9) or at least the amino acid sequence from amino acids No. 1 to 110 of Fig. 3 (SEQ ID No. 9).
 - 11. A purified BMP-9 protein obtainable by the steps of
 - (a) culturing a cell transformed with a cDNA comprising the nucleotide sequence from nucleotides No. 124 to 453 as shown in Fig. 3 (SEQ ID No. 8); and
 - (b) recovering and purifying from said culture medium a protein comprising the amino acid sequence from amino acids No. 1 to 110 as shown in Fig. 3 (SEQ ID No. 9).
 - 12. A purified BMP-9 protein obtainable by the steps of
 - (a) culturing a cell transformed with a cDNA comprising the nucleotide sequence from nucleotides No. 124 to 453 as shown in Fig. 3 (SEQ ID No. 8); and
 - (b) recovering from said culture medium a protein comprising an amino acid sequence from amino acids No. 8 to 110 as shown in Fig. 3 (SEQ ID No. 9).
- 13. A pharmaceutical composition comprising an effective amount of a protein according to any one of claims 7 to 12, optionally in conjunction with a pharmaceutically acceptable vehicle.
 - **14.** The composition of claim 13, further comprising a matrix for supporting said composition and providing a surface for bone and/or cartilage growth.
 - 15. The composition of claim 14 wherein said matrix comprises a material which is hydroxyapatite, collagen, polylactic acid or tricalcium phosphate.

- 16. The pharmaceutical composition of any one of claims 13 to 15 for wound healing, tissue repair, inducing bone growth or inducing cartilage growth.
- 17. Use of a protein according to any one of claims 7 to 12 for preparing a pharmaceutical composition for inducing bone formation, cartilage formation, treatment of wounds or tissue repair.
 - 18. A method for the preparation of a DNA sequence encoding a protein having the biological activity of a BMP-9 protein of inducing the formation of cartilage and/or bone which sequence is
 - (a) the DNA sequence from nucleotides 124 to 453 of SEQ ID No. 8; or
 - (b) the DNA sequence from nucleotides 145 to 453 of SEQ ID No. 8; or
 - (c) a DNA sequence which differs from the DNA sequence of (a) or (b) due to degeneracies of the genetic code;
 - (d) an allelic variant of the sequence of (a) or (b); or
 - (e) a DNA sequence hybridizing under stringent conditions to the sequences of (a) or (b),

said method comprising the following steps:

- (i) plating a human genomic library and preparing duplicate nitrocellulose replicas;
- (ii) hybridizing one set of the duplicate nitrocellulose replicas with the labeled oligonucleotide

#1: CTATGAGTGTAAAGGGGGTTGCTTCTTCCCATTGGCTGAT

and the other set with the labeled oligonucleotide

#2: GTGCCAACCCTCAAGTACCACTATGAGGGGATGAGTGTGG:

30 and

10

15

20

25

45

- (iii) isolating those clones which hybridize to both oligonucleotides and determining the sequence of their inserts.
- 19. A process for the manufacture of a composition according to claim 13, characterized in the use of the protein of any one of claims 7 to 12 as an essential constituent of said composition.

Patentansprüche

- DNA-Sequenz, die ein Protein mit der biologischen Aktivität der Induktion der Bildung von Knorpel und/oder Knochen eines BMP-9-Proteins codiert, wobei die Sequenz ist
 - (a) die DNA-Sequenz von Nucleotid 124 bis 453 von SEQ ID No. 8; oder
 - (b) die DNA-Sequenz von Nucleotid 145 bis 453 von SEQ ID No. 8; oder
 - (c) eine DNA-Sequenz, die sich von der DNA-Sequenz nach (a) oder (b) aufgrund der Degeneration des genetischen Codes unterscheidet; oder
 - (d) eine allelische Variante der Sequenz nach (a) oder (b); oder
 - (e) eine DNA-Sequenz, die unter stringenten Bedingungen mit den Sequenzen nach (a) oder (b) hybridisiert.
- Rekombinantes DNA-Molekūl, das eine DNA-Sequenz nach Anspruch 1 enthält.
 - 3. Rekombinantes DNA-Molekül nach Anspruch 2, wobei die DNA-Sequenz unter der Kontrolle von regulatorischen Elementen steht, die ihre Expression in einer gewünschten Wirtszelle erlauben.
- Wirtszelle, die das rekombinante DNA-Molekül nach Anspruch 2 oder 3 enthält.
 - 5. Wirtszelle nach Anspruch 4, die eine Bakterienzelle, eine Hefezelle oder eine Säugerzelle ist.

- 6. Verfahren zur Herstellung eines Proteins mit der biologischen Aktivität eines BMP-9-Proteins, umfassend die Züchtung einer Wirtszelle nach Anspruch 4 oder 5 unter Bedingungen, die für die Expression der DNA-Sequenz geeignet sind, und die Gewinnung des Proteins aus der Kultur.
- 5 7. Protein, das von der DNA-Sequenz nach Anspruch 1 codiert wird.
 - 8. Protein, das durch das Verfahren nach Anspruch 6 hergestellt wird.
 - 9. Protein mit der biologischen Aktivität eines BMP-9-Proteins, das eine der folgenden Aminosäuresequenzen umfaßt
 - (a) die Aminosäuresequenz von Aminosäure Nr. 8 bis 110, die in Fig. 3 (SEQ ID No. 9) dargestellt ist; oder
 - (b) die Aminosäuresequenz von Aminosäure Nr. 1 bis 110, die in Fig. 3 (SEQ ID No. 9) dargestellt ist.
 - 10. Protein mit der biologischen Aktivität eines BMP-9-Proteins, wobei das Protein ein Dimer ist, in dem jede Untereinheit mindestens die Aminosäuresequenz von Aminosäure Nr. 8 bis 110 von Fig. 3 (SEQ ID No. 9) oder mindestens die Aminosäuresequenz von Aminosäure Nr. 1 bis 110 von Fig. 3 (SEQ ID No. 9) umfaßt.
 - 11. Gereinigtes BMP-9-Protein, erhältlich durch die Schritte

10

15

30

40

50

55

- (a) Züchtung einer Zelle, die mit einer cDNA transformiert ist, die die Nucleotidsequenz von Nucleotid Nr. 124 bis 453 umfaßt, die in Fig. 3 (SEQ ID No. 8) gezeigt ist; und
 - (b) Gewinnung und Reinigung eines Proteins, das die Aminosäuresequenz von Aminosäure Nr. 1 bis 110 umfaßt, die in Fig. 3 (SEQ ID No. 9) gezeigt ist, aus dem Kulturmedium.
- 25 12. Gereinigtes BMP-9-Protein, erhältlich durch die Schritte
 - (a) Züchtung einer Zelle, die mit einer cDNA transformiert ist, die die Nucleotidsequenz von Nucleotid Nr. 124 bis 453 umfaßt, die in Fig. 3 (SEQ ID No. 8) gezeigt ist; und
 - (b) Gewinnung eines Proteins, das die Aminosäuresequenz von Aminosäure Nr. 8 bis 110 umfaßt, die in Fig. 3 (SEQ ID No. 9) gezeigt ist, aus dem Kulturmedium.
 - 13. Arzneimittel, das eine wirksame Menge eines Proteins nach einem der Ansprüche 7 bis 12 gegebenenfalls in verbindung mit einem pharmazeutisch verträglichen Träger, umfaßt.
- 14. Arzneimittel nach Anspruch 13, das weiter eine Matrix als Träger des Arzneimittels umfaßt und eine Oberfläche für Knochen- und/oder Knorpelwachstum bereitstellt.
 - 15. Arzneimittel nach Anspruch 14, wobei die Matrix ein Material umfaßt, das Hydroxyapatit, Collagen, Polymilchsäure oder Tricalciumphosphat ist.
 - **16.** Arzneimittel nach einem der Ansprüche 13 bis 15 zur Wundheilung, Gewebewiederherstellung, Induktion des Knochenwachstums oder Induktion des Knorpelwachstums.
- 17. Verwendung eines Proteins nach einem der Ansprüche 7 bis 12 zur Herstellung eines Arzneimittels zur Induktion der Knochenbildung oder der Knorpelbildung, zur Behandlung von Wunden oder zur Gewebewiederherstellung.
 - 18. Verfahren zur Herstellung einer DNA-Sequenz, die ein Protein mit der biologischen Aktivität der Induktion der Bildung von Knorpel und/oder Knochen eines BMP-9-Proteins codiert, wobei die Sequenz ist
 - (a) die DNA-Sequenz von Nucleotid 124 bis 453 von SEQ ID No. 8; oder
 - (b) die DNA-Sequenz von Nucleotid 145 bis 453 von SEQ ID No. 8; oder
 - (c) eine DNA-Sequenz, die sich von der DNA-Sequenz nach (a) oder (b) aufgrund der Degeneration des genetischen Codes unterscheidet; oder
 - (d) eine allelische Variante der Sequenz nach (a) oder (b); oder
 - (e) eine DNA-Sequenz, die unter stringenten Bedingungen mit den Sequenzen nach (a) oder (b) hybridisiert,

wobei das Verfahren die folgenden Schritte umfaßt:

- (i) Plattierung einer menschlichen genomischen Genbank und Herstellung von Nitrocellulose-Zweifachreplikas;
- (ii) Hybridisierung eines Satzes der Nitrocellulose-Zweifachreplikas mit dem markierten Oligonucleotid

5

#1: CTATGAGTGTAAAGGGGGTTGCTTCTCCCATTGGCTGAT

und des anderen Satzes mit dem markierten Oligonucleotid

#2: GTGCCAACCCTCAAGTACCACTATGAGGGGATGAGTGTGG;

15

und

(iii) Isolierung derjenigen Clone, die mit beiden Oligonucleotiden hybridisieren, und Bestimmung der Sequenz ihrer Insertionen.

20

30

35

19. Verfahren zur Herstellung eines Mittels nach Anspruch 13, dadurch gekennzeichnet, daß man ein Protein nach einem der Ansprüche 7 bis 12 als wesentlichen Bestandteil des Mittels verwendet.

25 Revendications

- Séquence d'ADN codant pour une protéine ayant l'activité biologique d'une protéine BMP-9 d'induire la formation de cartilage et d'os, laquelle séquence est
 - (a) la séquence d'ADN des nucléotides 124 à 453 de SEQ ID No. 8; ou
 - (b) la séquence d'ADN des nucléotides 145 à 453 de SEQ ID No. 8; ou
 - (c) une séquence d'ADN qui diffère de la séquence d'ADN de (a) ou (b) due aux dégénérescences du code génétique;
 - (d) une variante allèle de la séquence de (a) ou (b); ou
 - (e) une séquence d'ADN s'hybridant sous des conditions rigoureuses en les séquences de (a) ou (b).
- 2. Molécule d'ADN recombinant contenant une séquence d'ADN suivant la revendication 1.
- 3. Molécule d'ADN recombinant suivant la revendication 2, dans laquelle la séquence d'ADN est sous le contrôle d'éléments régulateurs permettant son expression dans une cellule hôte désirée.
 - 4. Cellule hôte contenant la molécule d'ADN recombinant suivant l'une ou l'autre des revendications 2 et 3.
- Cellule hôte suivant la revendication 4, qui est une cellule bactérienne, une cellule de levure ou une cellule mam mifère.
 - 6. Procédé de production d'une protéine ayant l'activité biologique d'une protéine BMP-9, comprenant la culture d'une cellule hôte suivant l'une ou l'autre des revendications 4 et 5 sous des conditions appropriées pour l'expression de la séquence d'ADN précitée et la récupération de ladite protéine de la culture.

- 7. Protéine codée par la séquence d'ADN de la revendication 1.
- 8. Protéine produite par le procédé de la revendication 6.
- 9. Protéine ayant l'activité biologique d'une protéine BMP-9 comprenant une des séquences d'acides aminés suivantes :
 - (a) la séquence d'acides aminés allant des acides aminés n° 8 à 110 telle que représentée à la figure 3 (SEQ

ID No. 9); ou

15

20

35

45

55

- (b) la séquence d'acides aminés allant des acides aminés n° 1 à 110 telle que représentée à la figure 3 (SEQ ID No. 9).
- 10. Protéine ayant l'activité biologique d'une protéine BPM-9, dans laquelle ladite protéine est un dimére dans lequel chaque sous-unité comprend au moins la séquence d'acides aminés allant des acides aminés n° 8 à 110 de la figure 3 (SEQ ID No. 9) ou au moins la séquence d'acides aminés allant des acides aminés n° 1 à 110 de la figure 3 (SEQ ID No. 9).
- 10 11. Protéine BMP-9 purifiée obtenable par les étapes suivantes :
 - (a) la culture d'une cellule transformée avec un ADNc comprenant la séquence nucléotidique allant des nucléotides n° 124 à n° 453 telle que représentée à la figure 3 (SEQ ID No. 8); et
 - (b) la récupération et la purification dudit milieu de culture d'une protéine comprenant la séquence d'acides aminés allant des acides aminés n° 1 à 110 telle que représentée à la figure 3 (SEQ ID No. 9).
 - 12. Protéine BMP-9 purifiée obtenable par les étapes suivantes :
 - (a) la culture d'une cellule transformée avec un ADNc comprenant la séquence nucléotidique allant des nucléotides n° 124 à n° 453 telle que représentée à la figure 3 (SEQ ID No. 8); et
 - (b) la récupération dudit milieu de culture d'une protéine comprenant une séquence d'acides aminés allant des acides aminés n° 8 à 110 telle que représentée à la figure 3 (SEQ ID No. 9).
- 13. Composition pharmaceutique comprenant une quantité efficace d'une protéine suivant l'une quelconque des revendications 7 à 12, éventuellement conjointement à un véhicule pharmaceutiquement acceptable.
 - **14.** Composition suivant la revendication 13, comprenant de plus une matrice pour supporter ladite composition et former une surface pour la croissance d'os et/ou de cartilage.
- 30 **15.** Composition suivant la revendication 14, dans laquelle ladite matrice comprend une matière qui est une hydroxyapatite, du collagène, de l'acide polylactique ou du phosphate tricalcique.
 - **16.** Composition pharmaceutique suivant l'une quelconque des revendications 13 à 15, pour cicatriser les blessures, réparer les tissus, induire une croissance osseuse ou induire la croissance de cartilage.
 - 17. Utilisation d'une protéine suivant l'une quelconque des revendications 7 à 12 pour préparer une composition pharmaceutique pour induire une formation osseuse, la formation de cartilage, le traitement de blessures ou la réparation de tissus.
- 18. Procédé de préparation d'une séquence d'ADN codant pour une protéine ayant l'activité biologique d'une protéine BMP-9 d'induire la formation de cartilage et/ou d'os, laquelle séquence est
 - (a) la séquence d'ADN des nucléotides 124 à 453 de SEQ ID No. 8; ou
 - (b) la séquence d'ADN des nucléotides 145 à 453 de SEQ ID No. 8; ou
 - (c) une séquence d'ADN qui diffère de la séquence d'ADN de (a) ou (b) due aux dégénérescences du code génétique;
 - (d) une variante allèle de la séquence de (a) ou (b); ou
 - (e) une séquence d'ADN s'hybridant sous des conditions rigoureuses en les séquences de (a) ou (b),
- 50 ledit procédé comprenant les étapes suivantes :
 - (i) l'étalement d'une bibliothèque génomique humaine et la préparation de répliques de nitrocellulose dupliquées;
 - (ii) l'hybridation d'une série des répliques de nitrocellulose dupliquées avec l'oligonucléotide marqué

n° 1: CTATGAGTGTAAAGGGGGTTGCTTCTTCCCATTGGCTGAT

et l'autre série avec l'oligonucléotide marqué

n° 2: GTGCCAACCCTCAAGTACCACTATGAGGGGATGAGTGTGG; et

5		(iii) l'isolement de ces clones qui s'hybrident aux deux oligonucléotides et la détermination de la séquence de leurs inserts.
10	19.	Procédé de fabrication d'une composition suivant la revendication 13, caractérisée par l'utilisation de la protéine suivant l'une quelconque des revendications 7 à 12 comme constituant essentiel de ladite composition.
15		
20		
25		
30		
35		
40		
45		
50		
<i>55</i>		

Figure 1A

<i>ር</i> አጥጥ እ	1 ממדמ	-	ኮጥ አ አ /	20 CTAT		ነ አ ጥጥ ል	30 CT 0	2 A A A T		10 G TT	ሶ ርጥጥ	50 cree		anne'	60 TGG (70 TGAGC	
CALIA		n in	T TUNA	OIAI	1001	WIII	.01 0	mni	1001	GII	CC11	0100	MG	JAMU.	100 (CAM	TONGC	
	8	0		90			100		11	.0		120)		130		140	
TTTTT	'AGTT	T GT	STCG	GAAG	CCT	TAAT	TA C	GGCI	CCAG	C TC	ATAG	TGGA	ATG	GCTA'	TAC 1	TAG?	TATTA	
	15	_		160			170		18	-		190			200		210	
GGATA			AGTA					rggta			TAGG		_	TTGA'		LATA	LATATG	
	22	_		230			240		2:	-		260	-		270		280	
ATTAG			ATTA				310	CGTCC		G GT 20	GTGT	GGAT 330		CATT	ATT 1	rgrri	MATAD	
ምእ አ ሮፕ	29 אמדייוויי	_	cmc s	300 TCT				20303			יייר א ייי			ביתים		TCC	350 SAATAG	
INNUI	36		GICA	370			380	JGAGA		90	IGNI	400		1110	410	11660	420	
GATTG		-	ТАТА	•				GATTO		-	TATT			GGGT		GAATO	AGGCA	
	43			440			450			50		470			480		490	
AATAG	TTTA	T CG	TTCA	TTTT	AAT	TCTC	AAG (GGGTT	TTTP	C TI	TATT'	GTTT	GTT.	AGTG	ATA :	PTGG:	rgagta	
	50	-		510			520			30		540	•		550		560	
GGCCA			ATAG			GAAT!			LAATO		TTAC			GATG			SAGGGC	
	5	70		58	0		590)		600		60	-		61	18		
	AAAGG										coo1	0000	·~ >.			- -		
TGA	AAAGG	CT C	CTTC	CCTC	:C C	AGGAC	AAA	A CCC	GAGC	AGG	GUCF	CCCC	M	S S	P		7 G	
													1.1	5	•	•		
	627			636			645			654			663			672		
GCC	TTC	CGG	GTG	GCC	CTG	CTC	CCG	CTG	TTC	CTG	CTG	GTC	TGT	GTC	ACA	CAG	CAG	
A	F	R	V	A	L	L	P	L	F	L	L	V	С	V	T	Q	Q	
	681			690			699			708			717			726		
776	CCG	CMC	2 2 2	220	===				mcc.	<u>~~</u>	~~~	CAN	2 200	<u> </u>	CAC	AGC	TCC	
AAG K	P	L	Q	N N	W	E	O	A	S	P	GGG	E	N	A	H	S	S	
K	r	ם	Q	14	"	L	Q	-	3	•	G	-	.,	••	••	_	_	
	735			744			753			762			771			780		
CTG	GGA	TTG	TCT	GGA	GCT	GGA	GAG	GAG	GGT	GTC	TTT	GAC	CTG		ATG	TTC	CTG	
L	G	L	S	G	Α	G	Ε	E	G	٧	F	D	L	Q	M	F	L	
													005					
	789			798			807			816			825			834		
CNC	AAC	NTC.	7 7 C		<u> </u>	TOTAL C	CTIA	666	AGC	<u> </u>	AAC	CTC	AGC	GGC	<u> </u>	\overline{CCC}	TCC	

Figure 1B

	843			852			861			870			879			888	
CAG Q			ACC T	AGA R	GCG A	GAG E	CCA P	CCC P	CAG Q		ĀTG M	ATC I	GAC D	TTG L	TAC Y	AAC N	ĀGĀ R
	897			906			915			924			933			942	
TAC Y	ACA T	ACG T	GAC D	AAA K	TCG S	TCT S	ACG T	CCT P	GCC A	TCC S	AAC N	ATC I	GTG V	CGG R	AGC S	TTC P	ĀGC S
	951			960			969			978			987			996	
GTG V	GAA E	GAT D	GCT A	ĀTĀ I	TCG S	ACA T	GCT A	GCC A	ACG T	GAG E	GAC D	TTC F	CCC P	TTT F	CAG Q	AAG K	CAC H
;	1005		:	1014		:	1023		;	1032			1041		:	1050	
ATC I	CTG L	ATC I	TTC F	AAC N	ATC I	TCC S	ATC I	CCG P	AGG R	CAC H	GAG E	CAG Q	ATC I	ACC T	AGG R	GCT A	GAG E
	1059			1068			1077			1086			1095			1104	
CTC L	CGA R	CTC L	TAT Y	GTC V	TCC S	TGC C	CAA Q	AAT N	GAT D	GTG V	GAC D	TCC S	ĀCT T	CAT H	GGG G	CTG L	GAA E
	1113			1122			1131			1140			1149			1158	
GGA G	AGC S	ATG M	GTC V	GTT V	TAT Y	GAT D	GTT V	CTG L	GAG E	GAC D	AGT S	GAG E	ACT T	TGG W	GAC D	CAG Q	GCC A
	1167			1176			1185			1194			1203			1212	
ACG T	GGG G	ACC T	AAG K	ACC T	TTC F	TTG L	GTA V	TCC S	ÇAG Q	GAC D	ATT I	CGG R	GAC D	GAA E	GGA G	TGG W	GAG E
	1221			1230			1239			1248			1257			1266	
ACT T	TTA L	GAĀ E	GTA V	TCG S	AGT S	GCC A	GTG V	AAG K	CGG R	TGG W	GTC V	AGG R	GCA A	GAC D	TCC	ACA T	ACA T
	1275	;		1284			1293	1		1302	!		1311			1320	
AAC N	ĀĀĀ K	AAT N	AAC K	CTC L	GAG E	GTG V	ACA T	GTG V	ÇAC Q	AGC S	CAC H	AGC R	GAG E	AGC S	TGT C	GAC D	ACA T
	1329	•		1338	3		1347	7		1356	5		1365	,		1374	
CTC L	G GAC	ĀTO	AG?	GTC V	CCI P	r CCA	G GG	TCC S	ĀĀJ K	AAC N	CTC L	CCC P	TTC F	<u>TT1</u> F	GTI V	GTC V	TTC F

Figure 1C

	1383		:	1392		:	1401		;	1410			1419		;	1428	
TCC S	AAT N	GAC D	CGC R	ĀGC S	AAT N	GGG G	ĀCC T	AAG K	GAG E	ACC T	ĀGĀ R	CTG L	GAG E	CTG L	AAG K	GAG E	atg M
	1437		:	1446		;	1455			1464			1473			1482	
ATC I	GGC G	CAT H	GAG E	CAG Q	GAG E	ACC T	ĀTG M	CTT L	GTG V	ĀĀG K	ACA T	GCC A	AAA K	AAT N	GCT A	TAC Y	CAG Q
	1491		:	1500		:	1509			1518			1527			1536	
GTG V	GCA A	GGT G	<u>GAG</u> E	AGC S	CAA Q	GAG E	GAG E	GAG E	GGT G	CTA L	GAT D	GGĀ G	TAC	ACA T	GCT A	GTG V	GGA G
	1545		;	1554		:	1563			1572			1581			1590	
CCA P	CTT L	TTA L	GCT A	AGA R	AGG R	AAG K	R	AGC S 319)	ACC T	GGA G	GCC A	AGC S	AGC S	Н	C	CAG Q	AAG K
	1599			1608			1617	319)		1626			1635	(,	326)	1644	
ACT T	TCT S	CTC L	AGG R	GTG V	AAC N	TTT F	GAG E	GAC D	ATC I	GGC G	TGG W	GAC D	AGC S	TGG W	ATC I	ĀTT I	GCA A
	1653			1662			1671			1680			1689			1698	
CCC P	AAG K	GAR E	TAT Y	GAC D	GCC A	TAT Y	GAG E	TGT C	AAA K	GGG G	GGT G	TGC C	TTC F	TTC F	CCA P	TTG L	GCT A
	1707			1716			1725			1734			1743			1752	
GAT D	GAC D	GTG V	ACA T	CCC P	ACC T	AAA K	CAT H	GCC A		GTG V		ACC T	CTG L	GTG V	CAT H	CTC L	GAG E
	1761			1770			1779			1788			1797			1806	
TIC F	CCC P	ACA T	AAG K	GTG V	GGC G	ĀĀĀ K	GCC A	TGC C		GTT V	CCC P	ACC T	AAA K	CTG L	ĀGT S	CCC P	ĀTC I
	1815			1824			1833			1842			1851			1860	
TCC	ĀTC I	CTC L	TAC Y	AAG K	GAT D	GAC D	ATG M	GGG G		CCA P	ACC T	CTC L	AAG K	TĀC Y	CAC H	TAT Y	GAG E
	1869			1878			1887		_		1	903		19	13		1923
GG G	ATG M	AGT S	:. <u>G</u> _C	GCT	GAG E	TGT C	GGG G	C	AGG R 428)		TCCC	TGC	AGCC.	ACCC.	AG G	GTGG	GATA

Figure 1D

1933	1943	1953	1963	1973	1983	1993
CAGGACATGG	AAGAGGTTCT					TCSTCS CSTS
2003	2013	2023				
CCATCCTTGA	GAAGAAAAGG					
2073	2083	2093				
GACTGGGGTA	TGCGGGCCTG					
2143	2153	2163				
	TGGGTAGATG					
2213	2223	2233				
	AGAACTCTGC				CTTGGGAGTG	TGTCCTCAGG
2283	2293	2303				
	TTGCTGTTCC	TGTGCCTCAA	GCTCCCAGCT	GACTCTCCTG	TGGCTCATAG	GACTGAATGG
2353	2363	2373				
GGTGAGGAAG	AGCCTGATGC	CCTCTGGCAA	TCAGAGCCCG	AAGGACTTCA	AAACATCTGG	ACAACTCTCA
242	3 243	3 244	13			
TTGACTGAT	G CTCCAACAT	AATTTTTAA	A AGAG			

Figure 2

7 (AGTGTG	GGTG	60 CTA	GAAG		50 CGGC	SAGCO	40 SC G		GAAG	30 AGG	AGGG.	GGG	20 AGGA	GAGG	-	1 GAGG	CTCTA
140 CTCCCC	TTGT	130 PAGC	'ATCI		120 GCTG	CTCCG		11 SCTG	CGCC	.00 AGA		GAG	90 ACGC	AGGG	C TG	80 CGAG	GCATO
210 TCACTO	AGGI	200 CCC	CTC		190 CCCG	CAGTO		18 AGCG	CTGC	.70 AGC	_	TAT	160 Aagc	GTCC		150 Gatt	GATGO
280 CATTC	GAGO	270 TATG	ACC1		260 TGCA	CTAC		25 CGAG	CTGG	40 CTG		GGA	230 CCCA	GGTC		220 GTTC	CAACO
350 AGATTGO	TCA	340 TTGT	AGTI		OEE ADDT	SCCT1		32 PCCC	AGCT	10 TGC		GCA	300 CAAC	CGAG		290 GCCA	GTAGI
	r cci	ATT			40 AGAC	GTCA	90 TC :		CCTI	380 TATG	TTAT		370 SACTO	CATGO		36 CAAGA	CTGT
																417 AAC Asn	
CAG Gln	ATT Ile	GAG Glu	GCC Ala	507 GTC Val	AAA Lys	AAA Lys	AAG Lys	GGG Gly	492 ACG Thr	GAG Glu	CCT Pro	ATA Ile	TTG Leu	477 AGT Ser	GCT Ala	CAT His	AGC Ser
																CAC His	
627 AAG Lys	AGC Ser	CCT Pro	CAG Gln	CCG Pro	612 CGC Arg	CGC Arg	CGC Arg	CTG Leu	GGG Gly	597 TTT Phe	ATG MET	CAG Gln	CTG Leu	CTT Leu	582 ACA Thr	GCG Ala	GAG Glu
GAG Glu	GGG Gly	TCT Ser	672 CAG Gln	CTT Leu	CGG Arg	TAC Tyr	CTT Leu	657 GAT Asp	CGG Arg	ATG MET	TAC Tyr	GAC Asp	642 CCG Pro	ATT Ile	GTC Val	GCC Ala	AGT Ser
GCC Ala	732 CCG Pro	CGC Arg	GAG Glu	CCT Pro	TAT Tyr	717 GAG Glu	CTI Leu	GGT Gly	ACT Thr	AGC	702 CAC His	ATC Ile	CAG Gln	GAG Glu	GAA Glu	687 GAG Glu	GAG Glu
ATC Ile	AAC Asn	GAG Glu	CTG Leu	777 CAT His	GAA Glu	GAA Glu	CAC His	CAC	762 TTC Phe	AGC Ser	AGG Arg	GTG Val	ACC Thr	747 AAC Asn	GCC Ala	CGG Arg	AGC Ser
ATC Ile	AGC	837 AGC Ser	CTC Leu	AAC Asn	TTT Phe	CTC	822 TTC Phe	CGT Arg	TTT Phe	GC1	TCT	807 AAC Asn	GAA Glu	AGT Ser	ACC Thr	GGG	792 CCA Pro

Figure 2A

CCT GAG Pro Glu																
GAC CAG Asp Glr																
957 ATG AAG MET Lys	CCC													CTA		
ACG AGA	CTG	GTC Val	CAC His	CAC His	AAT Asn	GTG	1032 ACA Thr	CGG Arg	TGG Trp	GAA Glu	ACT	TTT Phe	GAT Asp	GTG Val	AGC Ser	CCT Pro
1062 GCG GTG Ala Val			TGG					CAG					CTA			
GTG ACT					ACT					GGC					ATT	
CGA TC		CCT					AAT					CGG				
122 ACC TT Thr Ph	r GGC	CAT His	GAT Asp	GGC	1242 CGG Arg	GGC Gly	CAT His	GCC Ala	TTG	ACC Thr	CGA Arg	CGC Arg	CGG Arg	AGG	GCC Ala	AAG Lys
CGT AG Arg Se	CCT					CAG					AAG					
1332(3 CGC CA Arg Hi	TCG		TAT					GAT					GAC			
GCC CC Ala Pr	1392 A CCA o Pro	GGC	TAC	CAG Gln	GCC Ala	1407 TTC Phe	TAC	TGC Cys	CAT His	GGG	1422 GAC Asp	TGC Cys	CCC Pro	TTT Phe	CCA	CTG Leu
GCT GA Ala As	C CAC p His	CTC Leu	1452 AAC Asn	TCA	ACC Thr	AAC Asn	CAT His	1467 GCC Ala	ATT	GTG Val	CAG Gln	ACC	1482 CTG Leu	GTC	AAT Asn	TCT
149 GTC AA	7 T TCC n Ser	: AGT : Ser	ATC	CCC Pro	1512 AAA Lys	GCC	TGT Cys	TGT Cys	GTG Val	1527 CCC Pro	ACT	GAA Glu	CTG Leu	AGT	1542 GCC Ala	ATC

Figure 2B

TCC ATG CTG TAC CTG GAT GAG TAT GAT AAG GTG GTA CTG AAA AAT TAT CAG GAG Ser MET Leu Tyr Leu Asp Glu Tyr Asp Lys Val Val Leu Lys Asn Tyr Gln Glu 1602 1617 (408) 1636 1646 1656 ATG GTA GTA GAG GGA TGT GGG TGC CGC TGAGATCAGG CAGTCCTTGA GGATAGACAG MET Val Val Glu Gly Cys Gly Cys Arg ACAGACTGCT TCCTTATAGC TGGACTTTTA TTTAAAAAAA AAAAAAAAA AATGGAAAAA ATCCCTAAAC ATTCACCTTG ACCTTATTTA TGACTTTACG TGCAAATGTT TTGACCATAT TGATCATATA TTTTGACAAA

ATATATTAT AACTACGTAT TAAAAGAAAA AAATAAAATG AGTCATTATT TTAAAAAAAA AAAAAAAACT

CTAGAGTCGA CGGAATTC

Figure 3

	Thr			TGC												48
CAG Sln -25	GTG Val	AGA Arg	GCA Ala	GTC Val	ACG Thr -20	AGG Arg	AGG Arg	ACA Thr	CGG Arg	ATG Met -15	GCG Ala	CAC His	GTG Val	GCT Ala	GCG Ala -10	96
GGG Gly	TCG Ser	ACT Thr	TTA Leu	GCC Ala -5	AGG Arg	CGG Arg	AAA Lys	AGG Arg	AGC Ser 1	GCC Ala	GGG Gly	GCT Ala	GGC Gly 5	AGC Ser	CAC His	144
TGT Cye	CAA Gln	AAG Lys 10	ACC Thr	TCC Ser	CTG Leu	CGG Arg	GTA Val 15	AAC Asn	TTC Phe	GAG Glu	GAC Asp	ATC Ile 20	GGC Gly	TGG Trp	GAC Asp	192
AGC Ser	TGG Trp 25	ATC Ile	ATT Ile	GCA Ala	CCC Pro	AAG Lys 30	GAG Glu	TAT Tyr	GAA Glu	GCC Ala	TAC Tyr 35	GAG Glu	TGT Cys	AAG Lys	GGC Gly	240
GGC Gly 40	TGC Cys	TTC Phe	TTC Phe	CCC Pro	TTG Leu 45	GCT Ala	GAC Asp	GAT Asp	GTG Val	ACG Thr 50	Pro	ACG Thr	AAA Lys	CAC His	GCT Ala 55	286
ATC Ile	GTG Val	CAG Gln	ACC Thr	CTG Leu 60	Val	CAT His	CTC	AAG Lys	TTC Phe 65	Pro	ACA Thr	AAG Lys	GTG Val	GGC Gly 70	AAG Lys	336
GCC Ala	TGC Cys	TGI Cys	GTG Val 75	Pro	ACC	Lys	CTG Leu	AGC Ser 80	Pro	ATC Ile	TCC Ser	GTC Val	CTC Leu 85	Tyr	AAG Lys	384
GAT Asp	GAC Asp	ATG Met	: Gly	GTG Val	Pro	ACC Thr	CTC Lev	Lys	TAC Tyr	CAT His	TAC Tyr	GAG Glu 100	ı Gly	ATG Met	AGC Ser	432
		Glu		GGG Gly			ſ	TATO	CTGC	CTG	CGGG					470